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# GENERIC REVISION AND SKELETAL MORPHOLOGY OF SOME CERIOPORID CYCLOSTOMES (BRYOZOA) 

By<br>Osborne Barr Nye, Jr.

1976

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# GENERIC REVISION AND SKELETAL MORPHOLOGY OF SOME CERIOPORID CYCLOSTOMES (BRYOZOA) 

Osborne Barr Nye, Jr.<br>Syracuse University

## ABSTRACT

Thirteen post-Paleozoic cerioporid (Bryozoa) genera including 14 species have been restudied utilizing internal characters. This approach applied to routine studies of Paleozoic tubular Bryozoa has produced relatively consistent taxonomic schemes. Earlier studies of cyclostomatous Bryozoa were based on a relatively few, primarily external characters. Variations of these characters generally reflect non-genetic factors. The discovery of many new internal characters in post-Paleozoic cyclostomes expands the basis from which new taxonomies can be constructed and evolutionary inferences made. Presumably as biological relationships of internal and external structures become known, estimates of genetic and non-genetic factors which control their variation will improve.

Genera were diagnosed on the basis of characters associated with zoarial growth patterns, microstructure of the zooecial wall, and occurrence of diaphragms. Brood chambers, which are primary zoarial structures in the cerioporids studied, are too poorly known at present to provide taxonomic characters in supra-specific categories.

Cerioporids studied have ramose, massive, or frondose zoaria. Ramose habit was produced by: (1) the formation of an axial endozone composed of nearly parallel growing, thin-walled zooecia which eventually bend radially and become thick-walled in the exozone; (2) essentially like (1) as modified by a spiral budding pattern; (3) like (1), but zooecia stop growing orally after emplacement of frontal walls bearing peristomes; (4) repetitive hemispheric extensions of the basal layer to form an axial support structure upon which zooecia are initially adnate; (5) repetitive overgrowth in which each growth phase is composed of radially directed zooecia; (6) parallel growth of autozooecia which open only at growing tips. Frondose habit is produced by bifoliate budding from a median layer. Massive habit is produced by radial growth of zooecia. Overgrowth and intrazoarial anastomosis of growing branches are important modifications of growth habit in some genera.

Basal, intermediate, and terminal diaphragms; and simple external walls with restricted apertures can be identified in cerioporids. They can be distinguished on their position within the zooecium, direction in which laminae flex when merging with the zooecial wall, occurrence of pseudopores, and occurrence of peristomes. Basal, and perhaps intermediate, diaphragms formed floors to living chambers; terminal diaphragms presumably functioned as protective cover-plates to zooids in degenerative phases; simple external walls may have functioned as protective cover-plates by restricting the skeletal aperture to a small opening (peristome), through which feeding organs (the lophophore) had access to sea water. Basal diaphragms were secreted by membranes on the oral side of the diaphragm. Intermediate, terminal, and simple external walls were secreted by membranes on their aboral sides. The secretion of intermediate, terminal, and simple external walls is related to the connection of interzooidal tissue through interzooidal pores. Increased circulation through interzooidal pores, not possessed by most Paleozoic Bryozoa, may provide an adaptive advantage to most post-Paleozoic Bryozoa.

Observations of zooecial wall structure in cerioporids supports the "double wall" mode of growth model proposed by Borg (1926b, 1933) and expanded by Boardman and Cheetham (1969). In cerioporids, two major kinds of laminar structure can be distinguished. In one group, laminae arch orally convex. Four subgroups are distinguished on the basis of: (1) continuity of laminae across the zooecial boundary zone, (2) occurrence of subgranular calcite, and (3) occurrence of thick zooecial linings. In the second group, laminae intersect the axis of oral growth at less than $90^{\circ}$. In one subgroup, laminae are linear to slightly curved; in a second subgroup, laminae recurve aborally to form a broad arch in the outer cortex. The last subgroup occurs in a Bathonien species, thus extending the known occurrence of orally acute lamination.

## ACKNOWLEDGMENTS

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## ABBREVIATIONS FOR REPOSITORIES

| USNM | National Museu mof Natural History (formerly United <br>  <br>  <br> States National Museum), Smithsonian Institution, <br> Washington, D. C. |
| :--- | :--- |
| WM | British Museum (Natural History), London, Great Bri- <br> tain |

MNHN Institute de Paléontology, Muséum National d’Histoire Naturelle, Paris, France
NMW Naturhistorisches Museum, Vienna, Austria
UB Institut Palaontologie, Universität, Bonn, Federal Republic of Germany
ANSP Academy of Natural Sciences, Philadelphia, Pa., U.S.A.

## INTRODUCTION

Fossil genera of Cyclostome Bryozoa have been known since 1826 when Goldfuss erected the genus Ceriopora. Since that time, numerous cyclostome genera and species have been named, particularly in the works of Michelin (1841-1848), Haime (1854), von Hagenow (1851), d’Orbigny (1849b, 1854), Gregory (1896, 1902, 1909), and Canu and Bassler (1920, 1922, 1926). Knowledge of living cyclostomes has been increased by the efforts of Barrois (1877), Busk (1879), Waters (1879), Harmer (1890, 1893, 1897, 1899), Robertson (1903, 1910a, b), and Borg (1926a, 1933). The abundance of named species and genera and the length of time that they have been known suggests that cyclostome bryozoans should be, at present, a well-known group taxonomically. Yet this is not the case. Since the beginning of this century, cyclostomes have largely been relegated to the backwaters of taxonomic research. With the exception of Borg's investigations, fundamental understanding of cyclostomes has not advanced since about the turn of this century.

The major obstacle to the investigation of cyclostomes has been the lack of study techniques. In the past, most taxonomic studies were based on a few arbitrarily chosen external characters. Taken
singly or together, these characters were generally non-diagnostic by virtue of: a) their ubiquity throughout the cerioporids, e.g., "zooecial tubes cylindrical to prismatic", b) their ambiguity, e.g., "zooecial tubes long", or c) their having so much intertaxon variability as to be virtually useless. Definitions of taxa were unreliable and have not served to define or distinguish taxa. Illustration of the external characters of types has failed to provide sufficient documentation at the specific or generic level. Furthermore, largely because of homeomorphy, external characters are poor data from which to infer evolutionary relationships. As a result, existing taxonomic frameworks are inconsistent and largely unuseable. Thus, cyclostomes have been virtually ignored in geologic or biologic investigations which depend upon taxonomic information as basic data.

This study is an attempt to find new characters that will provide the data for the construction of a new taxonomic framework. One of the finest collections of fossil cyclostomes in the world is housed in the National Museum of Natural History. Numerous cyclostome species were thin-sectioned under the direction of R. S. Boardman during the summer of 1966. Preliminary examination of these sections indicated that cyclostomes have at least as many internal characters as Paleozoic Stenolaemata.

Species with relatively large or "stony" zoaria were easily thinsectioned by techniques in general use. Many of these species were referable to the Cerioporina, and the most recent comprehensive treatment of cerioporid genera was given by Bassler (1953). Therefore, the genera selected for this initial study were those assigned by Bassler to the Cerioporina as valid names or synonyms.

## TAXONOMIC BASIS AND PROCEDURE APPROACH

The major goals of this revision are two-part.The first is nomenclatural: to determine the validity of generic and specific names and to document types, primarily through photographic illustrations. Types are the objective fixtures of nomenclature and must form the nucleus of any revisionary taxonomic investigation.

Validation of generic names was facilitated by the large collection of literature on bryozoans collected by R. S. Bassler, later R. S. Boardman and A. H. Cheetham, and by the large general col-
lections of zoological literature in the National Museum of Natural History. Objective documentation of genera was approached through the location and redescription of the primary types of type species. When authoritative evidence indicated that the primary types were destroyed or lost from known repositories, generic names were retained only if secondary specimens could be assigned with confidence to the type species. This was necessary because most concepts based on external characters generally do not serve to define or distinguish cerioporid taxa. In each instance where concepts were based solely on examination of secondary specimens, the reasons for their use are discussed.

Internal characters are well known in Paleozoic tubular Bryozoa and provide the basis of internally consistent taxonomic concepts. It is reasonable to expect that the same approach should yield similar results when applied to the study of cyclostome bryozoans.

The second goal has been to formulate generic and specific concepts based primarily on skeletal structures, and to interpret skeletal structures biologically. In this first stage of revision, numerous internal structures were recognized. Choice of characters associated with certain structures does not imply inferences of phylogenetic importance but does expand the known phenotypic basis from which evolutionary inferences can be made. The concepts, if internally consistent, should provide the empirical data for second-level, more theoretical, studies, including the construction of taxonomies based on inferences of evolutionary linkage.

Construction of phylogenetic classifications implies knowledge of variation in genotypes through time. Estimates of genetic variation improve as nongenetic factors are excluded. In paleontology, variation in genotype is inferred from morphologic, primarily skeletal, characters. Boardman, Cheetham, and Cook (1969) have identified and discussed extragenetic elements which influence mode of growth in Bryozoa. These elements are ontogeny of zooids, astogeny, polymorphism, and microenvironment. Variation in these elements can be recognized in single colonies. Moreover, each colony is made up of numerous zooids and all zooids are virtually identical in genotype. Thus, investigators of colonial organisms have a powerful tool for calibration of extra-genetic sources of phenetic variation.

Taxonomic concepts, to be useful phylogenetically, should be based on characters which reflect genetic variability. Concepts developed here are, admittedly, preliminary because only the types are adequately prepared for study. Species descriptions are based on few specimens, and all but one genus are based on the examination of type species only. None the less, these concepts are not invalid; they are simply imprecise. A great deal of information can be derived from a few, or even single, specimens. Types have special bearing on nomenclature, but no special bearing on concepts. They are simply members of a population and, in terms of that population, bear no more and no less information than any other individual.

Concepts based on single specimens pose a special problem because concepts nominally imply knowledge of interspecimen variability. Herein, two species are presently known from lectotypes only. Because the specimens showed states of many characters assumed to have importance in other species, and because estimates of nongenetic variability can be made even from single zoaria, these specimens were fully described.

## GENERA INCLUDED

Of the approximately 50 cerioporid generic names listed by Bassler (1953), 17 are listed in Table 1 and represent progress to date on the generic revision of the group. Of the 19 names, four are objective synonyms, one is a subjective synonym, and one genus, Dysnoetopora, has been reassigned to the Cheilostomata (Voigt, 1971). The 13 remaining genera show relatively great variation in mode of growth and wall structure. In the future, it may be necessary to remove two of them (Corymbopora and Haploecia) from the cerioporids. Reassignment is not made here because all genera are compatible with Borg's double-wall concept and are not referable to the other existing double-walled groups, the hornerids, or lichenoporids. Many genera remain to be examined. Erection of new taxa at this stage is premature and could only serve to confuse rather than clarify.

## SYNONYMIES

The synonymies prepared here are objective in scope. They list those works which bear on the validity of names or documentation of types. Inclusion of non-objective references bears on taxonomic concepts, and in cerioporids, morphologic concepts as presently un-
derstood here must be based to a large extent on internal characters. Earlier investigators have based concepts on the relatively few external characters. Thus, published descriptions and illustrations are not sufficient for evaluation.

Relatively complete synonymies for names proposed prior to about 1900 are listed by Gregory (1896, 1899, 1909).

## GENERIC DIAGNOSES

Generic diagnoses, excepting that for Haploecia Gregory, are based on the type species. Information concerning specimens actually examined in this study is summarized in Table 1.

Characters (or character groups) believed to be useful at the generic level are:

1) Zoarial growth patterns, including the occurrence of polymorphism.
2) Microstructure of the zooecial wall.
3) Occurrence of diaphragms, and simple external walls.

In order to maintain consistency, characters based on structures observed in relatively few genera were excluded from generic diagnoses but were included in species descriptions. Brood chambers, for example, are striking morphological structures which are easy to identify and often have characteristic shapes. As such, various authors have considered them as important taxonomic characters at nearly all subordinal ranks (e.g., Canu and Bassler, 1920). In this study, brood chambers were observed in only five genera, and possibly a sixth (large primary chambers were observed in Ceriopora Goldfuss, but other structural characteristics typical of brood chambers were not observed). In four genera, the brood chambers were abundant and many occurred in each specimen. In the remaining genera, brood chambers were few; in Parleiosoecia Canu and Bassler, only three brood chambers were seen in 30 specimens. Brood chambers, therefore, were not included in generic diagnoses.

It is hoped that future investigations will clarify the occurrence and taxonomic importance of these structures.

## TECHNIQUES

When this investigation was begun, standard thin-section and peel techniques, as modified by R. S. Boardman and associates at
TABLE 1
GENERIC NAMES OF SOME POST-PALEOZOIC CERIOPORID BRYOZOA

|  | $\begin{aligned} & \text { Type Species } \\ & \text { \# Designation } \end{aligned}$ | Species Studied <br> *Type species | Type Specimen Designation | Materia <br> Syntypes | died Secondary specimens |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Ceriocava | SD Gregory, 1895 | $\begin{aligned} & \text { *Millepora } \\ & \quad \text { corymbosa } \end{aligned}$ | None | Probably destroyed | 10 Topotypes |
| Ceriopora | SD Gregory, 1895 | *C. micropora | Lectotype - here | Lectotype only | - - - |
| Corymbopora | M | *C. menardi | None | Probably destroyed | 11 Topotypes |
| Corymbosa d'Orbigny, 1854: Objective synonym of Corymbopora Michelin, 1846 |  |  |  |  |  |
| Coscinoecia | OD | *C. radiata | Lectotype - here | Lectotype only; others lost | - - - |
| \# SD - Subsequent designation <br> OD - Original designation <br> M - Monotypy |  |  |  |  |  |

GENERIC NAMES OF SOME POST-PALEOZOIC CERIOPORID BRYOZOA

|  | $\begin{aligned} & \text { Type Species } \\ & \text { \# Designation } \end{aligned}$ | Species Studied <br> *Type species | Type Specimen Designation | Mate <br> Syntypes | died Secondary specimens |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Dendroecia | OD | *Ceriocava muliilamellosa |  |  |  |
|  | Subjective synonym of Haploecia |  |  |  |  |
| Diplocava | OD | *D. incondita | Lectotype - here | Lectotype, 5 paralectotypes | 10 topotypes |
| Ditaxia | $\begin{gathered} \text { SD d'Orbigny, } \\ 1854 \end{gathered}$ | *Ceriopora anomalopora | Lectotype - here | Lectotype only | 8 topotypes |
| Dysnoetopora Canu and Bassler, 1920: Reassigned to the Cheilostomata (see Voight, 1971) |  |  |  |  |  |
| Haplocia | OD | *Millepora straminea | Lectotype Gregory, 1894 | Lectotype, 1 paralectotype | - - - |
|  |  | Ceriocava multilamellosa | Lectotype - here | Lectotype, 5 paralectotypes | 9 topotypes |

[^0]GENERIC NAMES OF SOME POST-PALEOZOIC CERIOPORID BRYOZOA

|  | $\begin{aligned} & \text { Type Species } \\ & \# \text { Designation } \end{aligned}$ | Species Studied <br> *Type species | Type Specimen <br> Designation | Mat <br> Syntypes | Studied Secondary specimens |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Hetcropora | SD Gregory, 1896 | *Ceriopora cryptopora | Lectotype - here | Lectotype, 5 paralectotypes | 11 topotypes |
| Lciosoccia | OD | *Multicrescis parvicella | Lectotype - here | Lectotype only | - - - |
| Parlciosoccia | OD | ${ }^{*}$ P. jacksonica | Lectotype - here | Lectotype, 2 paralectotypes | 31 topotypes |
| Polytaxia Hamm, 1881: Objective synonym of Ditaxia von Hagenow, 1851 |  |  |  |  |  |
| Reptonodicava | SD Bassler, 1935 | $\begin{gathered} \text { *Ceriopora } \\ \text { globosa } \end{gathered}$ | None | Probably destroyed | 12 topotypes |
| Spiroclausa d'Orbigny, 1854: Objective synonym of Zonopora d'Orbigny, 1849 |  |  |  |  |  |
| Tetrocycloccia | OD | *T. dichotoma | None | - - | 3 topotypes |
| Tretocycloccia Canu and Bassler, 1920: Objective synonym of Tetrocycloecia Canu, 1918 |  |  |  |  |  |
| Zonopora | OD | *Ceriopora spiralis | Lectotype - here | Lectotype only | 19 topotypes |
| $\begin{aligned} & \text { \# SD-Subsequent designation } \\ & \text { OD-Original designation } \\ & \text { M-Monotypy } \end{aligned}$ |  |  |  |  |  |

the National Museum of Natural History, were used (Boardman and Utgaard, 1964; Merida and Boardman, 1967). At that time, poorly indurated fossil cyclostomes and non-indurated Recent specimens were vacuum impregnated with polyester resins. In the course of this investigation, modifications of these techniques were made (Nye, Dean, and Hines, 1972). Essentially, these amounted to the utilization of epoxies for impregnation and mounting, and included fine polishing procedures of cut and ground faces. These modifications resulted in improved resolution of internal structures, the ability to section hard and soft parts together in Recent specimens, and the ability to make thin sections when desired (to approximately 5 microns).

## BIOMETRICS

Numerous characters were measured. A listing of these characters is given in Table 2. Phrases describing particular measurements are not always brief. Therefore, it was necessary to use abbreviations in statistical summaries of measurements found in each species description. The abbreviations are listed in Table 2.

Measurements of micro-dimensions were made directly through the microscope using an ocular micrometer. Projection techniques, which are faster, were attempted initially, but had to be abandoned because projected images of many specimens lacked sufficient contrast.

Commonly, more zooecia are available for measurement in tangential sections than it was feasible to measure, so a method of selection was necessary. Non-random methods of selection introduce bias and place constraints upon parametric statistics. Two random methods of selection were designed and are described below:

1) The microscope stage used could be moved parallel to two directions at right angles. A scale on the stage, calibrated to .1 mm , indicated the distance in each direction. The section to be measured was positioned, and the coordinates of the corners of a four-sided polygon which enclosed most of the section were noted. These coordinates were transferred to graph paper and a grid was constructed.

## TABLE 2

KEY TO ABBREVIATIONS USED IN STATISTICAL SUMMARIES

Zoarial
$\mathrm{Zr}-\mathrm{Ht}$
$\mathrm{Zr}-\mathrm{Wth}$
$\mathrm{Br}-\mathrm{CsSn}-\mathrm{MxDn}$
$\mathrm{PrBr}-\mathrm{CsSn}-\mathrm{MxDn}$
$\mathrm{Ov}-\mathrm{Th}$
AxCh-CsSn-MxDn

Bsilyy-Th
$\mathrm{BrCpm}-\mathrm{MxDn}$
$\mathrm{BrCpm}-\mathrm{MnDn}$
Zooecial
ZcCh-CsSn-MxDn
ZcCh-CsSn-NMxDn
Zc-LgnSn-Dph
ZcCh-CsSn-MxDn
$\overline{Z c C h}-\mathrm{CsSn}-\mathrm{NMxDn}$
$\mathrm{CdZcWl}-\mathrm{Th}$
ZdPr-Cn/ZcCsSn
ZdPr-MnDr
CnlZcCh-CsSn-MxDn
ZwWILn-Th
ZcSp-Cn/ZcCsSn
ZcCh-CsSn-LgnDn
ZcCh-CsSn-TrvDn

Simple External Wall
SEW-Th
SEW-Pst-CsSn-MxDn
SEW-Psdp-CsSn-MxDn

Zoarial Height
Zoarial Width
Branch - Cross Section - Maximum dimension
Primary Branch - Cross Section - Maximum dimension
(intrazoarial) Overgrowth - Thickness
Axial Chamber - Cross Section - Maximum Dimension
Basal Layer - Thickness
Branch Capitulum - Maximum Dimension
Branch Capitulum - Minimum Dimension

Zooecial Chamber - Cross Section - Maximum Dimension
Zooecial Chamber - Cross Section - Normal (to) Maximum Dimension
Zooecium - Longitudinal Section - Depth
Zooecial Chamber - Cross Section - Maximum Dimension
Zooecial Chamber - Cross Section - Normal (to) Maximum Dimension
Compound Zooecial Wall - Thickness
Interzooidal Pore - Count/Zooecial Cross Section
Interzooidal Pore - Minimum Diameter
Central Zooecial Chamber - Cross Section - Maximum Dimension
Zooecial Wall Lining - Thickness
(intra) Zooecial Spines - Count/Zooecial Cross Section
Zooecial Chamber - Cross Section - Longitudinal Dimension
Zooecial Chamber - Cross Section - Transverse Dimension

Simple External Wall - Thickness
Peristome in Simple External Wall - Cross Section Maximum Dimension
Pseudopore in Simple External Wall - Cross Section - Maximum Dimension

Diaphragm
TrlD-Th
TrlD-Psdp-CsSn-MxDn
IntD-Th
IntD-Intrl
IntD-DncApt
BsID-Th
BsID-Intvl

Terminal Diaphragm - Thickness
Pseudopore in Terminal Diaphragm - Cross Section - Maximum Dimension

Intermediate Diaphragm - Thickness
Intermediate Diaphragm - Interval
Intermediate Diaphragm - Distance (from) Aperture
Basal Diaphragm - Thickness
Basal Diaphragm - Interval

Brood Chamber

BrCh-Lth
BrCh-Wth
$\mathrm{BrCh}-$ Dth
$\mathrm{BrChFl}-\mathrm{Th}$
BcChRf-Th
BrChPsdp-Dr
Zoarial Position
NO
Ex
En
LgnSn
TngSn
TrvSn
Statistics

## OR

X
S
CV
N
NZc
NZr

Brood Chamber - Length
Brood Chamber - Width
Brood Chamber - Depth
Brood Chamber Floor - Thickness
Brood Chamber Roof - Thickness
Pseudopores (in) Brood Chamber Roof - Diameter

Not observed
Exozone
Endozone
Longitudinal Section
Tangential Section
Transverse Section

Observed Range
Mean
Standard Deviation
Coefficient (of) Variation
Number (of observations)
Number (of) Zooecia
Number (of) Zoaria

Each locus on the grid had an x and y coordinate. The number of zooecia to be measured was selected; then coordinates were chosen from a table of random numbers. The slide was positioned with respect to these coordinates, and the zooecium nearest the center of the field was measured. This method was time-consuming, as each tangential section required the construction of a new grid. Also, if zooecia were small and the zooecial wall thick, the coordinates were imprecise. This method was used to select zooecial characters in Reptonodicava globosa (Michelin) but was abandoned later in favor of the second method.
2) A photograph of the section was made and zooecia were numbered directly onto the photograph. Then numbers were selected from a table of random numbers. The zooecia so chosen were measured directly through the microscope. This method is fast; a polaroid $4 \times 5$ camera back was used, and prints were available within seconds. The method is precise, as well; if measurements are suspect, the zooecium can be found and dimensions checked.

Some measurements of zooecial characters are illustrated graphically for each species except Corymbopora menardi Michelin. The dimension normal to the longest dimension of the zooecial chamber was chosen because it should not be influenced by the
angular relation between the plane of the section and the zooecial growth axis. Also included are histograms of the ratio of major zooecial dimensions, compound zooecial wall thickness, and a cumulative curve for interzooidal pore counts.

Estimates of the arithmetic mean (X), the standard deviation (S), and coefficient of variation (CV) are not given for counts of interzooidal pores per zooecial cross-section. These counts do not meet the basic assumptions required for the use of parametric statistics; most importantly, when plotted, they do not approximate a normal distribution. The counts are summarized in cumulative curves given for each species.

## SKELETAL MORPHOLOGY

## ZOOECIAL WALL STRUCTURE

## MICROSTRUCTURE

Since the major studies by Ulrich commencing in the 1880's, wall structure has been considered an important taxonomic character in studies of Paleozoic stenolaemates. Nicholson was probably the first to make oriented thin-sections and observe skeletal microstructures in cyclostomes. He recognized and figured the laminar structure in the zooecial wall of Recent cerioporids (1880, p. 335; text-fig. 2, p. 336), Bleicher (1894, pp. 99-100, pl. 1, figs. 1, 3; pl. 2) prepared thin-sections and illustrated laminar structure in the zooecial wall of an encrusting tubuliporid cyclostome. Later investigators have misunderstood, or virtually ignored, microstructure.

In cerioporids, calcareous zooecial walls between adjacent zooecia are compound because they are grown from both sides. Therefore, in most genera, zooecial boundaries cannot be precisely defined because they lie within broad, tangentially-amalgamate zones. Laminae are sometimes arched continuously across the zone, or the zone may be composed of light-colored, subgranular, skeletal material which is nearly homogeneous in appearance. The continuity of calcareous tissue across the zooecial boundary zone suggests that the depositing epithelium passed continuously over the rims of adjacent zooecia. A membrane that included an outer cuticle covers the zoarium (observed by Borg, and probably Waters and Busk in Recent cerioporids, and by Harmer in Recent lichenoporids). The
outer membrane (gymnocyst of Borg) protects the inner depositing epithelium and probably aids in the transfer of nutrients around the actively growing apertural rims. Narrow, well-defined zooecial boundary zones can be seen in only a few genera. In these genera, the laminae of adjacent zooecia meet at relatively low angles.

Wall structure is not homogeneous throughout a zooecium. Zooecial walls are commonly homogeneous to subgranular, sometimes vaguely laminate in the thin-walled endozone and inner exozone portions. Thin zooecial linings are commonly present throughout. These are generally composed of dark-colored, longitudinally parallel laminae. Zooecia which bud from basal layers often have thick zooecial linings at the proximal tip of the zooecium and along the recumbent zooecial wall (Pl. 39, fig. 5).

Borg illustrated linear structures in the calcareous walls of Recent cerioporid species (1933, text-figs. 11, 15, 16, 17; pl. 7, figs. 5, 6). He referred to these, however, as fibers (1933, p. 337) and believed that they were organic, unspecified (e.g., see 1926a, p. 196), or chitinous (1926b, p. 585).

Recently, an integrative model of zooecial wall growth in Stenolaemata was presented by Boardman and Towe (1966, p. 20). The model was more fully developed by Boardman and Cheetham (1969, p. 211, text-fig. 2, p. 210). A similar approach has been used by Tavener-Smith (1969) and by Brood (1970a). This model integrates Borg's observations of the membranous portions of the body wall and Boardman and Towe's observations of microstructure and ultrastructures of the calcareous wall. Three-dimensional laminar configuration is the principal key to the understanding of skeletal morphology. Provided that the primary lamination is preserved, one can interpret structural relationships, sequence of events, and the location of the depositing epithelium (Boardman and Cheetham, 1969, p. 210). This model provides the basis for an understanding of the growth of zooecial walls in cerioporid bryozoans.

In the outer exozone of cerioporid genera, two major kinds of laminar microstructure can be distinguished. In one group, laminae arch orally convex, intersecting the orally directed axis of growth at $90^{\circ}$ or more (Text-fig. $1 \mathrm{~A}-\mathrm{D}$ ), and are orally oblique (Boardman and Cheetham, 1969, p. 211). In well-preserved specimens, laminae are continuous across the zooecial boundary zones (Text-fig. $1 \mathrm{~A}, \mathrm{~B}$ )


Text-figure 1 A-F. Diagrammatic profiles of compound zooecial walls in the outer exozone portion of cerioporid cyclostomes. Solid lines with arrows indicate inferred position of depositing portion of inner membrane responsible for last episode of cortex growth. Dashed lines in cortex indicate indistinct lamination; solid lines in cortex indicate distinct lamination; cross-hatching indicates subgranular to homogeneous calcareous tissue. In A-D, growth surfaces parallel lamination, and each lamina probably represents a single growth episode. In E and F, the growth surface parallels the depositing epithelia, but cuts across lamination; laminae probably grew by edgewise growth. Zooecial linings are included only in D. Linings may be deposited as sheetlike increments, or by edgewise growth.
or merge indistinctly with granular or homogeneous calcite in the outer cortex (Text-fig. 1 C, D). Each lamina is inferred to have been a simple growth surface which paralleled the depositing epithelium (Boardman and Cheetham, 1969, text-fig. 2A, p. 210).

In the second group, laminae intersect the axis of oral growth at less than $90^{\circ}$ (Text-fig. $1 \mathrm{E}, \mathrm{F}$ ) and are orally acute. Laminae meet with an angular relationship along the zooecial boundary zone producing an integrate appearance in tangential section (Pl. 18, fig. 3). Boardman and Cheetham (1969, p. 211, text-fig. 2B, p. 210) showed that laminae in this configuration were not parallel to the depositing epithelium and thus do not constitute single-event growth surfaces. Rather, growth is simultaneous along many laminae by deposition of calcareous crystals on the leading edge of each lamination.

Laminae within cerioporid zooecial walls extend aborally for only short distances and are not continuous with diaphragms. Most depositional activity, therefore, takes place at, and near, the apertural rim, and the deposition of diaphragms cannot be correlated with depositional events in the compound zooecial wall. Conversely, the zooecial walls of many trepostomes are composed of laminae which can be traced long distances aborally from the aperture. Often these laminae are continuous structurally with diaphragms, and form single diaphragm-wall units (Boardman, 1969, p. 27, text-fig. 8, textfig. 10, p. 31). In these, the membrane lining the entire living chamber apparently acted as a single depositional unit.

In cerioporids, microstructural subgroups can be distinguished. These are described below.

## Orally oblique lamination. -

Type 1. Laminae are broadly curved, arching continuously across the zooecial boundary zone (Text-fig. 1A). In tangential view, the zooecial walls are broadly amalgamate. This pattern has been observed in Ceriopora (Pl. 8, fig. 1d; Pl. 9, fig. 1a) Heteropora (PI. 35, fig. 2a) and Leiosoecia; and in the walls between adjacent small polymorphs in Ditaxia and Parleiosoecia (Pl. 40, fig. 1f).

Type 2. Laminae are broadly curved, arching continuously across the zooecial boundary zone. Laminated calcite alternates longitudinally with light-colored, heterogeneous to homogeneous calcite
(Text-fig. 1B). A better understanding of the light-colored calcite will necessitate investigation by electron microscopy. This tissue is inferred to be primary because it parallels well-preserved, laminated structures.

The light-colored calcite forms longitudinally discontinuous pluglike bodies in Coscinoecia (Pl. 14, fig. 1f) which are lapped by laminated calcite giving an acanthopore-like appearance in tangential section (Pl. 14, figs. 1g, h). These are not presently interpreted as acanthopores because the bodies are longitudinally discontinuous and because they lack structures typical of acanthopores in Paleozoic stenolaemates.

Type 2 is intergradational to some extent with Type 1 , but is most clearly distinguished in Coscinoecia (Text-fig. 1B, Pl. 14, fig. 1 e ; Pl. 15, fig. 1 g ).

Type 3. Laminae are nearly linear in profile (Text-fig. 10, Pl. 6, figs. 1, 3a) or arched (Pl. 5, figs. 1b, 1c). Laminae are distinct and closely spaced in the inner cortex. The outer cortex is lightcolored and homogeneous to indistinctly laminate (Pl. 5, figs. 1a, b, c; Pl. 6, figs. 1, 3a, 3b); laminae are sometimes seen to arch continuously across the zooecial boundary zone ( Pl . 5, fig. 1b). The poorly laminated appearance of the outer cortex is not simply an optical effect resulting from the angle of intersection between laminae and the plane of the section, because it was observed in longitudinal, transverse and tangential views. In addition, it was consistently observed in well-preserved specimens. Therefore, the appearance probably reflects some primary, but presently unknown, ultrastructure. This microstructure was observed in Ceriocava.

Type 4. The cortex is composed of light-colored subgranular to indistinctly laminated calcite. Laminae sometimes arch continuously across the zooecial boundary zone. In addition, the wall has a thick zooecial lining composed of dense, dark-colored, longitudinally directed, parallel to wavy laminae. The lining apparently thickens through ontogeny, and smooths over irregularities on the zooecial wall, such as spinose projections. This structure was observed in Haploecia and Zonopora and is illustrated in Text-fig. 1D, Pl. 24, fig. 1f; Pl. 26, fig. 1d; Pl. 30, figs. 1a, b; Pl. 31, fig. 2b; Pl. 47, fig. 1g; Pl. 48, figs. 1d, e, f; Pl. 49, figs. 1c, e; Pl. 50, figs. 1b, 2b, 2c.

Orally acute lamination. -
Type 5. In profile, laminae are linear to slightly curved, and commonly intersect the orally directed zooecial growth axis at about $45^{\circ}$ or less (Text-fig. 1E). Zooecial walls commonly are tangentially and longitudinally integrate in appearance. Thin zooecial linings composed of dark-colored, longitudinally directed laminae are commonly present. This microstructure was first illustrated by Borg (1933, text-fig. 11, p. 303; text-fig. 15, p. 322; text-fig. 16, p. 323 ; text-fig. 17, p. 329; text-fig. 20, p. 339; pl. 7, figs. 6, 7). Crystalline ultrastructures and mode of growth of this type were discussed by Boardman and Towe (1966, p. 20) and later, more fully, by Boardman and Cheetham (1969, p. 211; text-fig. 2B, p. 210; pl. 27, figs. 1a, 1b).

Type 6. Laminae initially extend from the zooecial growth axis at about $45^{\circ}$, then are broadly arched in the outer cortex. Zooecial walls are tangentially integrate in appearance. Zooecial linings are present or absent. This microstructure was observed in Diplocava and Reptonodicava and is illustrated in Text-fig. 1F; Pl. 16, figs. 1f, g, h; Pl. 17, fig. 6b; Pl. 18, figs. 2, 3; Pl. 19, fig. 1; Pl. 41, fig. 1 f .

## VARIATION IN THICKNESS

Zooecial walls of cerioporid bryozoans commonly show pronounced variation in thickness. In some genera this variation is cyclic, giving rise to annular thickenings ( Pl .4 , fig. 1e). More commonly, however, variation in thickness shows less regular patterns. Text-fig. 2 A-D illustrate the development of several different profiles in the outer exozonal zooecial walls of Coscinoecia radiata Canu and Lecointre. Moniliform profiles are enhanced by the occurrence of interzooidal pores but are not solely responsible for them. Interzooidal pores are nearly always located in thin-walled zones, but thin-walled zones are not always pierced by interzooidal pores.

A quantitative estimate of variation can be made from measurements of the width of compound zooecial walls in tangential section. The coefficients of variation generated from these measurements range from 27 in Ceriocava corymbosa (Lamouroux) to 55 for all polymorphs in Coscinoecia radiata Canu and Lecointre. The thickness of individual zooecial walls could not generally be measured because zooecial boundaries are not visible in thin-section. Further-
more, zooecial boundaries can be roughly approximated in longitudinal section by noting the orally directed crest of individual laminae (Text-fig. 2). Longitudinal lines connecting these points, the zooecial growth axes, are often significantly offset from the middle of the wall (Text-fig. 2C, D). Therefore, approximation of the zooecial wall thickness by halving the thickness of compound zooecial walls would often be inaccurate.

Qualitative estimates of variation in zooecial wall thickness can be made from profile views (Text-fig. 2 A-D). Species tend to show characteristic profiles and apparently have certain limits to variability. For instance, zooecial walls in Heteropora cryptopora (Goldfuss) are nearly parallel-sided (Pl. 34, fig. 1a) and become thicker orally in a regular manner. This contrasts with zooecial walls in Coscinoecia radiata Canu and Lecointre which commonly show great variation in thickness longitudinally (Text-fig. 2A, Pl. 14, fig. 1a).

There is some indication that this variation may be useful for delineation at supraspecific taxonomic ranks. For example, the zooecial walls near the branch tips of both Haploecia straminea (Phillips) and $H$. multilamellosa (Canu and Bassler) have irregular moniliform profiles. These become progressively more parallel-sided through the deposition of the zooecial lining.

The sample, however, appears too small to generalize from, and the important elements of profile shape, e.g., its relation to microstructure, are presently unknown. Therefore, estimates of profile shape are included here in species descriptions.

## DIAPHRAGMS AND SIMPLE EXTERNAL WALLS

## INTRODUCTION

Diaphragms are intrazooecial, calcareous partitions which extend transversely across zooecial tubes. They are found in the Paleozoic orders Trepostomata, Cryptostomata, and Cystoporata as well as the post-Paleozoic Cyclostomata. Haime (1854) was apparently the first to recognize diaphragms in the zooecial interiors of cyclostomes. Simple external walls, presently known only in cyclostomes, are discussed with diaphragms because they are much like terminal diaphragms in structure and position.


DISTAL G ROWTH OF
ZOARIUM

Text-figure 2 A-D. Incremental growth and longitudinal profiles of zooecial walls in the outer exozone of Coscinoecia radiata Canu and Lecointre. Solid lines indicate distinct laminae; broken lines indicate indistinct laminae; cross-checked patterns indicate subgranular to homogeneous-appearing tissue. Arrows indicate local zooecial growth directions. A. Profile is nearly symmetrical across the zooecial boundary zone, but shows large variation in thickness longitudinally. B. Profile is nearly symmetrical across the zooecial boundary zone and is almost parallel-sided. C. Profile is moderately variable in thickness longitudinally, and subsymmetrical across the zooecial boundary zone. Note longitudinal variation in direction of zooecial growth. D. Profile is subsymmetrical across zooecial boundary zone; note longitudinal variation in direction of growth.

Diaphragms in cerioporids generally are constructed of superposed calcareous laminae which are similar in appearance to the laminae of the zooecial lining. At the juncture with the zooecial wall, the diaphragms either flex orally or aborally. The orientation of this flexure can be used to infer the position of the depositing membrane. The flexure and depositing membrane are necessarily on the same side of the diaphragm if diaphragm laminae were deposited sequentially. In cerioporids, the flexed calcareous layers sometimes merge continuously with the zooecial lining. More commonly, the laminae adjoin the calcareous wall and extend for varying distances without merging, forming a structure referred to here as an abutment.

Three types of diaphragms can be identified in cerioporids: basal, intermediate, and terminal. Brief descriptions of these were given previously (Nye, 1970). The occurrence of diaphragms in 13 cerioporids is tabulated in Table 3. The diaphragms can be identified on the basis of:

1) the manner in which the diaphragms join the zooecial wall, 2) position with respect to the skeletal aperture, 3) presence or absence of pores.
Basal diaphragms are non-porous and generally thin, and with a few exceptions [e.g., in Haploecia straminea (Phillips), Plate 26, figure 1b], do not occur close to skeletal apertures. Basal diaphragms are primarily distinguished by the oral flexure of the diaphragm at the juncture with the zooecial wall (Pl. 5, fig. 2; Pl. 15, figs. 1b, f, i; Pl. 43, fig. 1). This oral flexure requires that the depositing epithelium was oral to the diaphragms (Text-fig. 3). Basal diaphragms of cerioporids and diaphragms of trepostomes are similar because they both flex orally at the juncture with the zooecial wall. In cerioporids, orally flexed calcareous tissue commonly forms thin abutments or merges obscurely with the zooecial lining. Thus, because basal diaphragms lack continuity with laminae in the cortex of the zooecial wall, deposition of basal diaphragms cannot be correlated with depositional events in the zooecial wall. This differs from some trepostomes in which Boardman (1960, p. 27, text-fig. 8) demonstrated that, ". . . a diaphragm plus the distally connected wall deposit are interpreted to form a unit of skeletal growth that was deposited at approximately the same time . . ."
TABLE 3
OCCURRENCE OF DIAPHRAGMS AND SIMPLE EXTERNAL WALLS IN SOME CERIOPORID GENERA

| Genus | Basal | Intermediate | Terminal | Simple external walls |
| :--- | :--- | :--- | :--- | :--- |
| Ceriocava | Numerous | Not observed | Common | Not observed |
| Ceriopora* | Not observed | Uncommon; just <br> subjacent to <br> overgrowth. | Not observed | Not observed |
| Corymbopora | Uncommon-Endo. | Not observed | Not observed | Not observed |
| Coscinoccia* | Numerous-Endo. | Common <br> zoarial overgrowth | Not observed | Not observed |
| Diplocava | Uncommon | Uncommon; close | Not observed | Common subjacent to |
| to aperture | Novergrowths |  |  |  |

*Based on a single specimen
OCCURRENCE OF DIAPHRAGMS AND SIMPLE EXTERNAL WALLS IN SOME CERIOPORID GENERA

| Genus | Basal | Intermediate | Terminal | Simple external walls |
| :--- | :--- | :--- | :--- | :--- |
| Heteropora | Uncommon | Common - subja- <br> cent to overgrowths <br> uncommon otherwise | Not observed | Not observed |
| Leiosoecia* | Not observed | Uncommon <br> exozone | Nouter | Not observed |

*Based on a single specimen


Text-figure 3. Diagrammatic profiles of terminal, intermediate and basal diaphragms in cerioporid bryozoans. All diaphragms were deposited from the inner membrane ( $=$ cryptocyst of Borg). The inner membrane is extended over the exterior side of the terminal diaphragms; this interpretation is consistent with Borg's observations of soft tissues (1933, text-fig. 2, p. 369). Arrows are emplaced at the approximate position of membranes which deposited the diaphragms.

Coscinoecia has a well-developed coaxial mode of growth. Basal diaphragms are numerous and closely spaced in the endozone and the zone of zooecial bending, but they were not observed in the exozone (Pl. 13, figs. 1d, g). This contrasts with most trepostomes in which diaphragms are commonly absent or infrequent in the endozone but numerous in the exozone (Boardman, 1960, p. 22).

Boardman (1960, p. 34) demonstrated that ". . . diaphragm counts and width of ephebic zones can be proportional to growth stages of zooecia considering the manner of skeletal growth." In the exozones of Ceriocava (Pl. 1, fig. 1 g ) and in Reptonodicava ( Pl . 44, fig. 2c), basal diaphragms are numerous and closely spaced. Ceriocava is robustly branching with ramose growth habit, and diaphragms are more numerous distally from growing tips. Reptonodicava has a massive growth habit in which zooecia essentially grow semiradially. Zooecia growing parallel to the major axis of distal growth are long, and have numerous diaphragms; zooecia growing at an angle to the major axis of zoarial growth are shorter and have fewer diaphragms. In Reptonodicava and presumably other bryozoans with similar growth habits, the number of diaphragms is both a function of ontogeny and a function of zooecial growth direction relative to the major axis of zoarial growth.

Intermediate diaphragms flex aborally at the juncture with the zooecial wall (Pl. 9, figs. 1a, b; Pl. 15, fig. 1a; Pl. 36, fig. 1g) and are the same in this regard as most terminal diaphragms. Intermediate diaphragms differ from terminal diaphragms in being nonporous. In addition, they are commonly thinner and are seldom observed at the skeletal aperture. Utgaard (1968b, pp. 1445-46, pl. 181, fig. 6) reported similar diaphragms in the ceramoporoid genera Ceramoporella Ulrich and Acanthoceramoporella Utgaard. Utgaard speculated that the diaphragms ". . . may be associated with a terminal phase of a zooid."

As noted by Utgaard (1968b, p. 1445), the aboral flexure of intermediate diaphragms requires that the soft tissues which deposited the laminae lay on the aboral side of the diaphragm (Textfig. 3). Soft tissues engaged in metabolic activities such as the deposition of calcareous tissue presumably require a supply of nutritive and respiratory substances. This, in turn, requires either a storage facility or a direct communication with tissues able to supply
these requirements. When the first calcareous lamina of the diaphragm is completed, a chamber is formed which is sealed off from the overlying zooecial cavity. Because of the lack of pores through the diaphragm, any significant transfer of metabolites would be eliminated. In cerioporids, however, soft tissues within these chambers presumably have access to nutrients via the zoarial communication system of interzooidal pores.

Terminal diaphragms (Borg, 1933, p. 290) are deposited at, or close to, the distal extremity of a single zooecium as zooecial cover plates. They are characterized by their position: aboral flexure, forming abutments (Pl. 5, fig. 1a; Pl. 6, fig. 3a; Pl. 40, fig. 1f) or merging with the zooecial lining (Pl. 50, figs. $2 \mathrm{~b}, \mathrm{c}$ ); and by the occurrence of pseudopores. Terminal diaphragms probably occur in most cyclostomes; they have been observed by several authors, including Busk (1859), Nicholson (1880), Waters (1884a), Pergens (1890), Robertson (1910), Canu and Bassler (1922, 1926), and Borg (1933, 1944).

The epithelial tissue which deposited the terminal diaphragms was on the aboral side of the diaphragms. This is so because diaphragms flex aborally at the juncture with the zooccial wall, and because laminae form U-shaped figures between pseudopores.

This interpretation of growths leads to the inference that diaphragms and pores were externally sealed by a cuticular layer. If so, the pores are pseudopores in the original sense of Borg and they probably functioned as cites of gas exchange.

In this respect, Borg (1944, pp. 76, 116) referred to the pores in terminal diaphragms as pseudopores. This contradicts his earlier opinions in two ways: (1) he believed (1933, pp. 277-8, 289, 356) that calcite was deposited on both sides of terminal diaphragms and (2) he believed (1933, p. 368, text-fig. 26, p. 369) that ectodermal and mesodermal tissue of the inner wall (cryptocyst) lined both sides of the diaphragm. It follows that Borg must have regarded any diaphragmal pores as true pores in his original sense until his 1944 opinion. It is unfortunate that no new anatomical evidence was cited in 1944 when he called the pores, pseudopores.

Terminal diaphragms are zooecial cover plates and probably serve to protect the soft tissues of the zoarial interior. In postPaleozoic cyclostomes, the breaching of a single zooecium can pro-
vide access to the entire zoarial interior via the interzooidal pores. Zoarial protection was first ascribed to terminal diaphragms by Waters (1884a, p. 403): "Now if each zooecium during its polypideless condition could be choked up by the mud deposited from the sea, then the whole colony might suffer". Harmer (1890) observed a damaged zooecium which was sealed by a terminal diaphragm. Later, a new polypide grew in the interior and the diaphragm was resorbed. Borg (1944, p. 116) also observed zooecia which were broken and then sealed by terminal diaphragms.

Zooids lacking feeding structures are often sealed by terminal diaphragms. Harmer (1890) observed that a ". . . zooecium which possesses a diaphragm contains a brown body, but no functional polypide. Here and there it will be noticed that a polypide-bud is being developed below the diaphragm. With the further development of this bud, the diaphragm is absorbed, the mouth of the zooecium again growing out into a long tube." In some dimorphic species, small polymorphs which probably did not house feeding zooids are commonly sealed by terminal diaphragms (Borg, 1933, pp. $289,368)$. Large polymorphs are generally open, and closure probably follows the degeneration of a feeding zooid. Perhaps external environmental stresses, e.g., seasonal variations, may induce closure. In this case, nearly all zooecia in the zoarium would probably be sealed, as is the case observed in some zoaria of Ceriocava corymbosa Lamouroux (PI. 1, figs. 1d, f). Closure would provide some protection to housed soft tissues. Presumably these tissues could survive periods of low nutrition until conditions improved, at which time feeding structures could be regenerated and the protective cover plates resorbed.

Simple external walls are apertural closures which are characterized by the presence of pseudopores. In Haploecia Gregory and Diplocava Canu and Bassler they are, in addition to pseudopores, pierced by a single, large peristome (Pl. 16, fig. 1g; Pl. 18, figs. 1-3; Pl. 28, fig. 1c). The peristome is the restricted skeletal aperture through which tentacles were probably protruded. The peristome often extends orally from the surface of the diaphragm ( Pl . 24, fig. 1f).

The simple external walls in Haploecia Gregory and Diplocava Canu and Bassler may be homologous to the structure in the Eleidae
called an "opercule" by D'Orbigny (1854, p. 606), or to the zooecial cover plates discussed by Levinsen (1912) in the eleid genus Melicerities d'Orbigny. The structures in Meliceritites, called opercula by Levinsen, are probably terminal diaphragms. Viskova (1968, text-fig. 16, p. 176) figured a thick section of a zooecial cover plate in Brachysoecia grandis Viskova which is probably a simple external wall. Canu and Bassler $(1922,1926)$ referred to the simple external wall as "facettes perforated by an aperture".

In Haploecia Gregory and Diplocava Canu and Bassler the structure of simple exterior walls resembles terminal diaphragms in that laminae commonly flex aborally at junctures with the zooecial walls to form abutments or merge with zooecial lining, and laminae form U-shaped figures between pores (Pl. 18, fig. 1; Pl. 26, fig. 1a). The wall is simple because it was grown from the interior side only and the skeletal evidence requires a depositing epithelium interior to the diaphragm. Then, as argued for terminal diaphragms, the exterior side of the diaphragm was lined by cuticle and the pores are pseudopores probably functioning as cites of gas exchange.

Simple external walls differ from terminal diaphragms in having restricted apertures commonly extended orally by peristomes. In addition, the emplacement of the simple wall marks the end of oral growth of compound zooecial walls and are thus considered as external calcareous walls. In many tubular bryozoans characterized by coaxial endozone-exozone mode of growth, zooecia continue to grow orally from the zone of flexure for some distance, and commonly intersect the zoarial surface at $90^{\circ}$. Boardman (1960, p. 34) showed that exozones characteristically become thinner in a gradient projected distally along a branch. Boardman used this relationship as an indicator of ontogeny in the interpretation of zoarial growth. The only apparent limit to the oral growth of an individual zooecium was the death of the depositing tissue (Boardman, 1960, p. 39).

In Haploecia, exozones do not show consistent change in thickness, apparently reaching their maximum thickness just proximal to the growing tip of the branch. Furthermore, zooecia never grow orally beyond the zone of zooecial flexure, and consequently intersect the zoarial surface at $40^{\circ}$ to $60^{\circ}$, never growing beyond a zone referable to the innermost exozone of many other species. Examination of unencrusted growing tips (Pl. 23, figs. 1c, d) indicates that ex-


Text-figure 4 A-E. Diagrammatic profile showing inferred development of simple external walls. Shaded areas are compound zooecial wall and simple external wall; parallel lines indicate zooecial lining and peristome wall; cross-hatched pattern is basal layer and zooecial walls of intrazoarial overgrowth; dark line labeled OM is outer, cuticle-lined membrane; dashed line is inner membrane; arrows indicate sites of active calcareous deposition by inner membrane; and ? indicates questionable extent of inner membrane.
ternal simple walls are emplaced coincidentally with the termination of calcification at the oral tips of the zooecial walls.

Boardman, Cheetham, and Cook (1969, explanation of textfigure 6) noted that continued growth of exozonal zooecia in a primary branch is sometimes arrested by superposition of an overgrowth, and that the full ontogenetic development of zooecia in the primary branch is never achieved. The thin exozones of Haploecia straminea might be considered to exemplify the above, if only those branches with extensive encrustations were observed. Several unencrusted branches of the type species, however, were sectioned and these also show no increase in diameter proximally.

There is evidence for the continued inhabitation of the zooecial cavity by soft tissues, and continued ontogenetic development after emplacement of the simple external wall and termination of orallydirected growth of the zooecial wall. The walls are thicker, and more often structurally complex in a gradient oriented proximally from the growing tip (Text-fig. 12). Also, zooecial linings are seen to be thicker in more proximally situated zooecia. Continued deposition of skeletal tissue requires the presence of depositing epidermis. In addition, peristomes probably were large enough to allow the protrusion of tentacles. It would seem possible that most of the zooecial chambers having access to the zoarial surface housed living, functional zooids.

## INTERZOOIDAL PORES

The term interzooidal pore is equivalent to the terms interzooecial pore, mural pore, communication pore, infundibular pore, septulae, and canaliculi as used by other authors. Interzooidal pores are canals through the calcareous compound walls between adjacent zooids which connect the body cavities of adjacent zooids. This usage is slightly modified from the definition given by Borg (1926a, p. 201).

Interzooidal pores are not equivalent in morphology or function to pseudopores. Pseudopores pass through calcareous single walls and are bounded externally by cuticle (Borg, 1926a, text-fig. 2, p. 193). Functionally, pseudopores are thought to allow restricted communication (i.e., of dissolved gases) to the exterior, whereas interzooidal pores allow communication between adjacent zooids.

The nearly ubiquitous presence of interzooidal pores in cyclostomes and their probable function was well known to several authors in the latter part of the last century. In 1879, Busk and Waters published separate observations on Recent cerioporid species in which they commented upon interzooidal pores. Busk (1879, p. 725) believed that the pores allowed ". . . the permeation of fluids throughout the entire zoarium". Nicholson (1880, p. 332) noted the importance of the findings of Busk and Waters, and (p. 421) compared the structure of the pores to mural pores in the Favositidae, noting that the interzooidal pores in cerioporids have " . . . definite walls and dilated extremities instead of being mere circumscribed deficiencies in the wall".

Examination of the microstructure of the zooecial walls in cyclostomes shows that the laminae lining the wall generally parallel the zooecial cavity until they reach the locus of the pore. Here the laminae do not stop abruptly, but deflect and contour the outline of the pore. Deflection of the laminae indicates that calcareous tissue was deposited by a secretory membrane which lined the pore; therefore, the pore is considered a primary structure.

Most Paleozoic bryozoans have nonporous walls. Interzooidal communication is limited to connections through zoarial tissues outside of the apertures (Boardman and Cheetham, 1969, p. 214). Ordovician and Silurian ceramoporids have communication pores (Utgaard, 1968a, b, 1969), but they do not appear to relate structurally to the interzooidal pores of post-Paleozoic cyclostomes. Most post-Paleozoic bryozoans have porous walls. Waters (1884b, pp. 6767) believed that interzooidal pores were complete homologues of the simplest of rosette plates among the cheilostomes. The constriction observed by Waters, however, is not a separate structure ( Pl . 5 , figs. 1c, 3; Pl. 6, figs. 2, 3b; Pl. 30, fig. 3; Pl. 51, figs. 2a-c), and interzooidal pores differ anatomically from the interzooidal communications in the Gymnolaemata (Borg, 1926a; Silén, 1944; Banta, 1969).

The existence of soft tissues lining the pores was first noted by Busk (1879, p. 725), and later confirmed by Harmer (1896) and Borg (1926a, pp. 201-202). Most authors believed that the interzooidal communication was open. Borg (1926a, p. 201), for example, wrote that, ". . . the zoids [sic] in the Cyclostomata have a much
more open communication with each other than is the case in the Cheilostomata and the Ctenostomata". Harmer (1893, p. 213), however, observed strands of funicular tissue which passed through the pores.

Protrusion of the polypide is affected through increase in turgor pressure acting upon the membranous sac (Borg, 1923, pp. 7-8). If the interzooidal pores are open, increase in turgor pressure should be transmitted throughout the zoarium. Clark (1964, p. 104) considered this problem and stated that, "The existence of restraints to polypide eversions is important since they permit the independent eversion and retraction of polypides which share a common hydrostatic skeleton". The restraints listed by Clark included the polypide retractors, relaxation of the vestibule dilator muscles, and contraction of the vestibular sphincter.

Preliminary examination of sections with both soft and hard tissues indicates that at least some pores are nearly completely filled by a single large cell (Pl. 51, figs. 2b, c). Sometimes the soft tissue appears to be connected with a tenuous network of connective tissue (Pl. 51, fig. 2c); other pores seen in the same section, however, appear to be devoid of large cells or connective tissue.

Interzooidal communication combined with the ability to secrete protective calcareous diaphragms, was probably one factor in the success of post-Paleozoic cyclostomes. In most Paleozoic bryozoans, the secretion of diaphragms within a zooecium formed a series of closed chambers. Living soft tissues were confined to a zone at the periphery of the zoarium, defined and underlain proximally by the last-formed diaphragm. This skeleton would seem to have provided only a supporting function analogous to that of a coralline calyx with a relatively small protective potential. In post-Paleozoic cyclostomes, however, living tissues capable of metabolic activities could be supported throughout the zoarial framework because of the communication system of interzooidal pores. This tissue may provide a temporary internal reservoir in time of stress when external conditions might be unfavorable to the existence of most feeding zooids. Under more favorable conditions, these underlying tissues might support the proliferation of new feeding zooids allowing the survival of the cyclostome colony. Thus, post-Paleozoic bryozoans, pro-
vided with interzooidal communications, have a flexibility in reacting to environmental changes not possessed by most Palcozoic bryozoans.

## ZOARIAL BROOD CHAMBERS

Occurrence and taxonomic importance of zoarial brood chambers have been discussed by Waters (1890), Harmer (1896), Canu ( $1898,1899,1918,1919$ a, b), Canu and Bassler (1930), and Borg (1926a, 1933, 1944). Borg (1933, pp. 267-9, pl. 2, figs. 1-3) demonstrated the presence of larvae in typical brood chambers of Recent cerioporid species.

In cerioporids, zoarial brood chambers are large skeletal cavities found in exozones (Pl. 1, fig. 1f; Pl. 3, figs. 1c, 3; Pl. 12, fig. 2c; Pl. 15, figs. 1c, e, f; Pl. 32, fig. 1f; Pl. 34, fig. 2b ). General structures are illustrated in Text-fig. 5. The chambers have more-or-less complete floors which are compound walls and which seal off most subjacent zooecia. The lateral walls are compound walls shared with adjacent zooecia. The chambers are commonly covered by a porous calcareous roof which is similar in appearance to the terminal diaphragms (Pl. 6, fig. 3c; Pl. 33, fig. 1; Pl. 40, figs. 1c, d, e). The nature of the membrane exterior to the brood chamber roof is presently unknown. Borg (1926a, p. 407) stated only that the calcified roof is " . . . a cryptocyst beneath the original thin gymnocyst"; thus, the pores through the roof are not called pseudopores. If it could be shown that pores are directly sealed by cuticle, then the pores are pseudopores in the sense of Borg. Following abandonment, brood chambers are commonly submerged beneath basal layers which advance from the compound lateral wall and from which new zooecia bud and grow (Pl. 3, fig. 1c; Pl. 32, fig. 1f; Pl. 33, fig. 1). Coscinoecia retains a continuous skeletal opening to the surface for some time after it is initially overgrown. In most other cerioporid genera, however, brood chambers are completely submerged.

In some genera, zooecia pass through the interior of the chamber, appearing like supporting columns. These intrachamber zooecia often have thin-walled septate partitions which extend laterally from them (Pl. 32, fig. 1f; Pl. 33, fig. 1; Pl. 51, fig. 1). The partitions commonly radiate away from the central open area. The intrachamber zooecia have thin compound walls. These walls were formed

Text-figure 5. Composite profile of abandoned brood chambers in cerioporid cyclostomes.
by back-to-back deposition from the zooidal membrane and the membrane lining the brood chamber. The septate walls are thinner, and were probably deposited by the brood chamber membrane only (Text-fig. 5, Pl. 32, fig. 1f; Pl. 33, fig. 1).

Borg (1933, pp. 269-70) believed that brood chambers were formed by the secondary resorption of the walls of zooecia adjacent to a fertile zooecium. If the chambers had arisen through resorption, then the floor and wall of the brood chamber would have been plastered over subjacent zooecial walls unconformably. Thus, microstructures of the zooecia and the brood chambers should be discontinuous and marked in thin-section by a sharp boundary zone. The brood chambers observed in this study, however, are primary structures. The floor and lateral walls of the brood chambers are compound walls and structurally continuous with subjacent or adjacent zooecial walls (Pl. 6, fig. 3c; Pl. 12, figs. 2d, e; Pl. 15, fig. 1c; Pl. 33, fig. 1 ; Pl. 40 , fig. 1d). In addition, pores are sometimes seen to connect brood chambers to adjacent zooecial chambers. These pores appear in all respects as true interzooidal pores formed as primary structures at the time the wall was deposited.

The compound growth of the partitions bounding the brood chamber indicates that the chamber should be considered as a zoarial structure rather than an inflation of a single fertile zooid (gonoecium). The zoarial membranes at apertures of numerous zooecia acted in concert to deposit the floor. In Coscinoecia, poorly-preserved microstructure suggests that the floor may have been formed by the progressive ringlike growth of a terminal zooecial closure (Pl. 15, fig. 1h). In some zooecia, intermediate diaphragms were secreted subjacent to the floor (Pl. 15, fig. 1h). In other genera, the floor was initially a single continuous closure and deposition continued on both sides.

## SYSTEMATIC DESCRIPTIONS

$$
\text { Order CYCLOSTOMATA Busk, } 1852
$$

Suborder CERIOPORINA von Hagenow, 1851
Genus CERIOCAVA d'Orbigny, 1854
Type species: Millepora corymbosa Lamouroux, 1821, p. 87, pl.

83, figs. 8, 9 by subsequent designation, Gregory (1896, p. 162).
1821. Pars Millepora Lamouroux, Exposition Méthodique des Genres de l'Ordre des Polypiers, des Zoophytes d'Ellis et Solander, p. 87, pl. 83, figs. 8, 9.
1849. Non Monticulipora d’Orbigny, Rev. et Mag. Zoologie, vol. 1, ser. 2, p. 503.
1854. Ceriocava d’Orbigny, Terrain Crétacé Bryozaires: Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 1016.
1896. Ceriocava d'Orbigny, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, p. 162.
1922. Ceriocava d'Orbigny, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 90 .
1935. Monticulipora d'Orbigny, Bassler, Fossilium Catalogus, I, Pars 67, Bryozoa, pp. 14, 69.
1953. Ceriocava d'Orbigny, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G70.
1953. Non Monticulipora d'Orbigny, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G70.

Tentative diagnosis. - Zoaria branching; branches having strongly differentiated coaxial endozone and exozone. Zoarial surface smooth to monticular.

In endozone, zooecial cross sections markedly small relative to those in exozone. Basal diaphragms widely spaced.

In exozone, zooecial walls indistinctly to distinctly laminate. Laminae commonly forming broadly rounded to $V$-shaped patterns pointing orally. Laminae merging obscurely with structurally indistinct tissue at zooecial boundary zone. Basal diaphragms more numerous and closely spaced than in endozone. Terminal diaphragms common.

Taxa included. - Only C. corymbosa Lamouroux, the type species, and $C$. multilamellosa Canu and Bassler were studied. C. multilamellosa was designated the type species of Dendroecia by Cotillon and Walter (1965) and is reassigned here to Haploccia because of the occurrence of peristomial diaphragms, mode of growth, and zooecial wall structure. The internal characters of other species assigned to Ceriocava are unknown to me.

Discussion. - D'Orbigny listed 13 species in Ceriocava with no indication as to which were typical. Gregory (1896, p. 162) designated C. corymbosa as the type species because it is the first recognizable species in D'Orbigny's list, and because it is the oldest and best known species.

Bassler (1934, p. 408) discovered that D'Orbigny (1854) had
designated Monticulipora "frustulosa" (Michelin) (a misspelling for the trivial name pustulosa Michelin, 1846) as the type species of Monticulipora (also Utgaard and Boardman, 1965, p. 112). Bassler (1935, p. 151; 1953, p. G70) listed Ceriopora pustulosa Michelin, 1846 (= Monticulipora "frustulosa" d’Orbigny, 1854) as synonymous with Millepora corymbosa Lamouroux, 1821 (the placement of Ceriopora pustulosa Michelin in synonymy under Millepora corymbosa Lamouroux is not here evaluated because of the lack of information concerning internal characters of Michelin's species). Unfortunately, Bassler (1935) also listed Ceriopora pustulosa as the type species of Ceriocava. This last action is illegal under provisions of the ICZN because it violates priority in view of Gregory's earlier (1896) designation of Millepora corymbosa Lamouroux as the type species. Secondly, Bassler's designation violates provisions concerning availability because Ceriopora pustulosa did not appear in D'Orbigny's original list of 13 species of Ceriocava, and deliberately so since he had already designated it as the type species of Monticulipora.

Because of his belief that Ceriopora pustulosa was the type species of both Monticulipora and Ceriocava, Bassler (1935, p. 151) cited Ceriocava as a junior synonym of Monticulipora. In 1953 (p. G70) and without explanation, Bassler reversed his earlier position and listed Monticulipora as an objective synonym of Ceriocava.

At the petition of Bassler and Duncan, the ICZN (1955, Opinion 443) designated Monticulipora mammulata d'Orbigny, 1850, as the type species of Monticulipora d'Orbigny, 1849. Thus at present, there is no justification for placing Monticulipora in synonymy with Ceriocava or vice-versa, with or without regard to the conspecificity of Ceriopora pustulosa Michelin and Ceriocava corymbosa Lamouroux.

D'Orbigny established the genus Ceriocava in order to distinguish ". . . tous les Ceriopora des auteurs ayant une seule couche de cellules et des ouvertures simples, représentant dans leur ensemble, une colonie rameuse" (1854, p. 1015). Gregory (1896, p. 163) emphasized the " . . . thick, irregularly branching habit . . ." and added information on some internal characters: "The axis of the zoarium consists of fine zooecia densely packed. The outer zone consists of zooecia which are usually reflexed and of much greater
diameter." Gregory illustrated a specimen of Corymbosa, the type species, with numerous diaphragms in the endozone (text-fig. 13, p. 164). Canu and Bassler (1922, pp. 90-2, text-fig. 20, p. 91) also described and illustrated thin sections of this species. The illustrations clearly show numerous diaphragms in the endozone, and an increase of zooecial diameters in the exozone relative to the endozone. One character not mentioned or illustrated is the occurrence of basal diaphragms in the exozone.

Ceriocava differs from Coscinoecia in having basal diaphragms in the exozone, and in having a V-shaped laminar structure and light-colored tissue in the outer exozone which often narrowly outlines the zooecial boundary zone. These genera differ in other characters such as details of brood chambers, architecture, and occurrence of mural spines; but these characters are tentatively considered diagnostic at the specific level.

Ceriocava differs from Reptonodicava in having branched zoaria with well-differentiated coaxial exozone-endozones, and in wall structure.

> Ceriocava corymbosa (Lamouroux), $1821 \quad$ Pl. 1, figs. 1a-h; Pl. 2, figs. 1a-h; Pl. 3, figs. 1a-c, 2, 3; Pl. 4, figs. 1a-e; Pl. 5, figs. 1a-c, 2, $3 ;$ Pl. 6, figs. 1, 2, 3a-c
1821. Millepora corymbosa Lamouroux, Exposition Méthodique des Genres de l'Ordre des Polypiers, des Zoophytes d'Ellis et Solander, p. 87, pl. 83, figs. 8, 9.
1854. Ceriocava corymbosa (Lamouroux), d'Orbigny, Terrain Crétacé Bryozoaires: Paléontologic Française Description des Animaux Invertébrés, vol. 5, p. 1016.
1896. Ceriocava corymbosa (Lamouroux), Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural Histry), the Jurassic Bryozoa, pp. 163, 164, text-fig. 13, p. 164.
1922. Ceriocava corymbosa (Lamouroux), Canu, and Bassler, U.S. Nat. Museum, Proc., vol. 61, p. 90, pl. 14, figs. 5, 6, 8; p. 91, fig. 20.
1965. Ceriocava corymbosa (Lamouroux), Cotillon, and Walter, Soc. Géol. France Bull., vol. 7, ser. 7, p. 935.

Type.-Lamouroux' specimens of C. corymbosa, originally stored in the Museum at the Université de Caen, were probably destroyed during the invasion of Normandy in World War II (fide Prof. L. Dangeard, Université de Caen). Lamouroux did not designate a holotype, and no specimen is known to have been designated as the lectotype.

Type locality and horizon. - "Terrain à polypiers environ de Caen" (Jurassic, Bathonien, Calvados, France).

Material studied. - Thin sections and acetate peels were made from the following topotypes: MNHN IP2-1, IP2-2, Bathonien, Ranville (Calvados), France; USNM 32164-1, $-2,-3$, Bathonien, St. Aubin (Calvados), France; USNM 32181-1, -2, -3; USNM 68941-1, -2, Bathonien, Ranville (Calvados), France. USNM 68941-1 was figured by Canu and Bassler, 1922, p. 14, fig. 8 and USNM 68941-2, pl. 14, fig. 6. Duplicate acetate peels are preserved in the National Museum of Natural History, Washington, D.C., and the author's collection.

Description. -
Growth habit. - Branches are robust, and have subcircular to elliptical cross sections. Monticules are ridgelike to pustulose. Intrazoarial overgrowths occur, but are generally local in extent.

The exozone and endozone intergrade in a broad zone of zooecial bending. Endozonal zooecia grow approximately parallel to the major branch axis for relatively long distances (Pl. 4, fig. 1e). Orally from the zooecial bend, zooecia are commonly rectilinear in growth and intersect the zoarial surface at approximately $90^{\circ}$.

Branches are massively intergrown in axillary zones distal to the bifurcation of branches (Pl. 2, fig. 1h; Pl. 4, fig. 1b). In the axillary zone, zooecia in each new branch are recurved distally as the distance separating the growth surfaces of each branch approximates 1 mm . Distally from this axillary zone of bending, the zooecial walls intersect obliquely. At the intersection the walls anastomose, forming a continuous compound wall against which zooecial chambers are pinched out (Pl. 4, fig. 1b).

Zones of irregular zooecia were occasionally observed. In one mode, zooecia are small in diameter, thin-walled, and irregular in growth direction (Pl. 3, figs. 1b, 2). In a second mode (Pl. 3, fig. 1a), the zooecia bud from a diaphragm-like structure, sealing off subjacent zooecial cavities. Superiacent zooecial walls are structurally continuous with the diaphragm-like structures.

Endozone - Zooecial walls show cyclic repetition of structure (Pl. 1, fig. 1d; Pl. 2, fig. 1h). In each cycle (Pl. 4, fig. 1e), individual zooecial walls are thin and straight to slightly undulatory for a rela-
tively long distance. Distally, marking the boundary of the cycle, zooecial walls are thickened annularly and form a zone parallel to the growing tip of the branch. Zooecial cross sections are small and elliptical to subelliptical (PI. 2, fig. 1f). Interzooidal pores were rarely observed. Zooecial walls are dark in color and indistinctly granular with a thin zooecial lining. Basal diaphragms (Pl. 5, fig. 2) are thin (less than .0016 mm ), dark in color and commonly convex aborally, and they flex orally to merge with the zooecial lining.

Exozone - Zooecial walls are generally symmetrical (less commonly slightly asymmetrical) in thickness across the zooecial boundary zone ( Pl .1 , fig. lg; Pl. 5, fig. 1c). Zooecial walls generally have distinctly moniliform profiles due, in part, to the occurrence of numerous, large and widely flared interzooidal pores ( Pl .5 , fig. 1c; Pl. 6, fig. 3b). Moniliform profiles are commonly oblate to elliptical near the zooecial bend (Pl. 3, fig. 1a), becoming alate to sagitate orally (Pl. 3, fig. 1c; Pl. 4, figs. 1c, d). Zooecial chambers are commonly subcircular in cross section. Mural spines are abundant (Pl. 5, fig. 3; Pl. 6, fig. 2) to nearly absent (Pl. 1, fig. 1h).

Diaphragms - Terminal diaphragms are numerous. The diaphragms are thick (Table 4, TrlD-Th) and are commonly seen to occur slightly aboral to the skeletal aperture (Pl. 1, figs. 1d, f; Pl. 2, figs. 1e, f; Pl. 5, fig. 1a; Pl. 6, fig. 3a). The oral surfaces of the diaphragms are generally planar, but the aboral surfaces are often uneven (Pl. 5, fig. 1a). The diaphragms sometimes adjoin the zooecial wall with slight flexure, or have relatively thick aborally flexed abutments (Pl. 5, fig. 1a; Pl. 6, fig. 3a). Diaphragms occasionally show slight flexure towards the aperture (Pl. 5, fig. 1a; proximal portion of top diaphragm).

Basal diaphragms are numerous and closely spaced (see Table 4, BaslD-Int; Pl. 1, fig. 1f; Pl. 2, fig. 1e; Pl. 3, fig. 1a). The diaphragms are thin (less than .0016 mm ) and slightly convex aborally, and they merge obscurely with the zooecial lining (Pl. 5, fig. 1c; Pl. 6, figs. 3a, b).

Brood chambers - Roughly oblate brood chambers were observed in the exozone (Pl. 1, fig. 1f; Pl. 2, fig. 1g; Pl. 3, fig. 1c). The chambers are large; measurements of the longest dimension parallel to the zoarial surface ( $\mathrm{BrCh}-\mathrm{Wth}$ ) and of the dimension
normal to the zoarial surface ( $\mathrm{BrCh}-\mathrm{Dth}$ ) are given in Table 4. The floor is complete but is pierced by occasional interzooidal pores ( Pl . 6, fig. 3c). The porous roof is thick (Table 4, BrChRf-Th). Abandoned brood chambers are submerged beneath a thin basal layer. Zooecia budding from the basal layer are either thick-walled with moniliform profiles and relatively straight in growth, or they are thin-walled, parallel-sided and irregular in growth (Pl. 3, fig. 1c).

Discussion. - Unfortunately, the external characters of Ceriocava corymbosa as figured by Lamouroux are not diagnostic. Later authors have referred to C. corymbosa, but none had redescribed or refigured the type specimens prior to their presumed destruction (see "Type" above). Under such circumstances, the assignment of specimens to Lamouroux' species and continued usage of the name is open to some question. The species concept, however, is reasonably consistent from author to author as seen in the descriptions and illustrations of Gregory (1896, pp. 164-5, text-fig. 13, p. 164), Canu and Bassler (1922, pp. 90-92, text-fig. 20, p. 91, pl. 14, figs. 1-8; 1929, p. 115, pl. 1, figs. 4, 5) and Cotillon and Walter (1965, p. 935). Furthermore, specimens examined by Gregory and Canu and Bassler were collected from the type locality, and the authors cited above considered internal characters in their diagnoses. For these reasons, and for the purpose of stability, the continuation of $C$. corymbosa (Lamouroux) seems justified.

Remarks concerning mode of growth. - Zoaria of C. corymbosa exhibit the typical coaxial exozone and endozone mode of growth of most large ramose Stenolaemata. Also, the periodic annular thickening of the zooecial walls in the endozone is essentially similar to cyclic growth phenomena demonstrated in other stenolaemates, such as in three living cerioporid species (Borg, 1933), two Devonian trepostome species (Boardman, 1960) and in the Upper Paleozoic trepostome, Tabulipora Young (Gautier, 1970). Intrazoarial overgrowths were observed only in localized areas, and do not appear to play a major role in zoarial increase. One specimen of $C$. corymbosa was found which completely encrusts a branch fragment of Haploecia straminea (Phillips).

The formation of an axillary zone of intergrown zooecia presumably functions to strengthen the colony. The recurvature and anastomosing growth of zooecia is not unique to C. corymbosa but

## TABLE 4

STATISTICAL SUMMARY OF MEASUREMENTS OF CERIOCAVA CORYMBOSA (LAMOUROUX)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} Z \mathrm{r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zoarial } \\ & \text { Br-CsSn-MxDn } \end{aligned}$ | 11.5 | 2.9 | 7.4 | 2.7 | 37 | 7 |  | 7 |  |
| $\begin{aligned} & \text { Zooecial-Exozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 38 | . 14 | . 28 | . 04 | 13 | 250 | 250 | 6 | 1 |
| Zch-CsSn-NMxDn | . 30 | . 11 | . 23 | . 03 | 13 | 250 | 250 | 6 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{Z \mathrm{Zch}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 2.1 | 1.0 | 1.2 | . 2 | 13 | 250 | 250 | 6 | 1 |
| CdZcWl-Th | . 14 | . 03 | . 08 | . 02 | 27 | 250 | 250 | 6 | 1 |
| ZdPr-Cn/ZcCsSn | 5.0 | 0 |  |  |  | 250 | 250 | 6 | 1 |
| ZcSp-Cn/ZcCsSn | 13.0 | 0 | 3.8 | 2.6 | 68 | 250 | 250 | 6 | 1 |
| ZdPr-MnDr | . 019 | . 011 | . 015 | . 002 | 15 | 25 | 5 | 1 | 5 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF CERIOCAVA CORYMBOSA (LAMOUROUX)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} \mathrm{Zr}_{\mathrm{r}}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zooecial - Endozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 13 | . 03 | . 09 | . 02 | 21 | 75 | 75 | 3 | 2 |
| Zch-CsSn-NMxDn | . 12 | . 02 | . 07 | . 02 | 25 | 75 | 75 | 3 | 2 |
| $\frac{Z c \mathrm{Ch}-\mathrm{CsSn}-\mathrm{MxDn}}{Z \mathrm{Cch}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 2.1 | 1.0 | 1.3 | . 2 | 18 | 75 | 75 | 3 | 2 |
| $\mathrm{CdZcWl-Th}$ | . 04 | . 01 | . 02 | . 01 | 38 | 75 | 75 | 3 | 2 |
| ZdPr-Cn/ZcCsSn | 2.0 | 0 |  |  |  | 75 | 75 | 3 | 2 |
| ZcWlLn-Th | . 003 | 0 |  |  |  | 50 | 50 | 2 | 3 |
| $\begin{aligned} & \text { Diaphragm - Exozone } \\ & \text { TrlD-Th } \end{aligned}$ | . 12 | . 02 | . 05 | . 02 | 46 | 21 |  | 3 | 6 |
| TrlDPr-CsSn-MxDn | . 025 | . 012 | . 017 | . 004 | 23 | 18 |  | 1 | 5 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF CERIOCAl'A CORYMBOSA (LAMOUROUX)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} Z_{\mathrm{r}}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Diaphragm-Exozone (con.) } \\ & \text { BsiD-Int } \end{aligned}$ | . 50 | . 08 | . 23 | . 08 | 34 | 75 | 15 | 3 | 4 |
| Brood Chamber BrCh-Wth | . 7 | . 5 |  |  |  | 2 |  | 1 | 5 |
| $\mathrm{Br}_{r} \mathrm{Ch}-\mathrm{Dth}$ | . 3 | . 3 |  |  |  | 2 |  | 1 | 5 |
| BrChFl-Th | . 40 | . 14 | . 27 | . 09 | 34 | 9 |  | 1 | 5 |
| BrChRf-Th | . 56 | . 41 | . 48 | . 05 | 10 | 11 |  | 1 | 5 |

[^1]TABLE 5
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN CERIOCAVA CORYMBOSA (LAMOUROUX)*

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |  |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: | :--- |
| Exozone <br> Regular | 1 |  |  |  |  |  |  |  |
| Sub | 142 | 38 | 17 |  |  |  |  |  |
| Irregular | 24 | 22 | 4 |  |  |  |  |  |
| Endozone |  | 14 | 3 | 3 |  |  |  |  |
| Regular |  |  |  |  |  |  |  |  |

*Estimates of exozone cross-sections were made from MNHN IP2-1 (50 zooecia), IP2-2 ( 100 zooecia), USNM 32181-1 (3 zooecia), USNM 32181-2 ( 25 zooecia), USNM 68941-2 ( 25 zooecia) and USNM 32164-2 ( 25 zooecia). Endozonal zooecial shapes were estimated from MNHN IP2-2 (25 zooecia), USNM 32164-2 ( 25 zooecia) and USNM 68941-2 ( 25 zooecia).
was observed at the oblique intersection of distally growing branches of Haploecia multilamellosa (Canu and Bassler) and Parleiosoecia jacksonica Canu and Bassler. R. S. Boardman (pers. comm. 1971) stated that zooecial recurvature is typical of trepostomes in the same situation.

The recurvature of zooecia suggests that the initial stimulus for change in growth habits is related to the proximity of zooecia growing towards each other from each new branch. Possibly the stimulus is provided by direct contact of soft tissues. Recurvature, however, begins when zooecia are separated by as much as 1 mm ; thus, if direct contact provides the stimulus, contact would be limited to the tips of tentacles.

The mode of anastomosis is similar to that seen in Parleiosoecia jacksonica in that zooecia growing towards each other tend to become recurved, and to approach the surface of contact obliquely. In both species, zooecia form compound walls of anastomosis with opposing zooecia, and individual zooecia are pinched out as growth continues. Anastomosis of zooecia, however, commonly occurs at the juncture of two distally growing branches in $P$. jacksonica rather than following the bifurcation of a single branch into two branches. Also in $P$. jacksonica, the zooecial walls of intersecting zooccia be-


Text-figure 6 A-D. Histograms and cumulative curve from three topotypes of Ceriocava corymbosa (Lamouroux). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.
come thin-walled and parallel-sided prior to merger. In C. corymbosa the zooecial walls remain thick-walled and have moniliform profiles up to the merger of intersecting zooecial walls. In addition, in C. corymbosa the new wall formed at the zone of intersection is moniliform in profile and thick-walled (Pl. 4, fig. 1b).

Irregularities in budding habit and zooecial appearance occur locally in the exozone in C. corymbosa. In one mode, new zooecia are budded from thick diaphragm-like structures which seal off subjacent zooecia (Pl. 3, fig. 1a). In a second mode, normally appearing zooecial walls terminate and are followed by irregularly oriented, thin-walled zooecia. Although the zooecial chambers appear to be continuous from normal to irregular zooecia, the zooecial walls are discontinuous and the walls of irregular zooecia are continuous with adjacent zooecial walls (Pl. 3, fig. 2). These irregularities are interpreted as a specialized zoarial response to disease or trauma by which zoarial growth may proceed. If so, this differs from other species in which zoarial repair is affected by the extension of a basal layer over dead portions of the zoarium, e.g., Reptonodicava (PI. 43, figs. 3a, b; Boardman, 1960, p. 39).

## Genus CERIOPORA Goldfuss, 1826

Type species: Ceriopora micropora Goldfuss, 1826, p. 33, pl. 10, figs. 4a, d, by subsequent designation, Gregory (1896, p. 195).
1826. Pars Ceriopora Goldfuss, Petrefacta Germaniae, vol. 1, p. 32.
1830. Pars Ceriopora Goldfuss, Blainville, Zoophytes: Dictionnaire de Science Naturelles, vol. 60, p. 378.
1834. Pars Ceriopora Goldfuss, Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 413.
1851. Pars Ceriopora Goldfuss, von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, p. 52.
1851. Pars Heteropora Blainville, von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, p. 52.
1896. Pars Ceriopora Goldfuss, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, p. 195.
1909. Pars Ceriopora Goldfuss, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 156.
1953. Ceriopora Goldfuss, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G57.
1953. Non Reptonodicava d'Orbigny, Bassler, Treatise on Invertebrate Paleontology: Part G, Bryozoa, p. G57.
Tentative diagnosis. - Zoaria massive with radial growth habit modified by development of major growth axis. Zoarial increase
in part of repetitive, but irregularly occurring, intrazoarial overgrowth. Endozone commonly restricted to portions of zooecia just adjacent to basal wall of intrazoardial overgrowths. In exozone, zooecial walls exhibiting variable repetition of thin-walled and thickwalled phases.

In thin-walled phase of exozone, zooecial wall granular to indistinctly laminate. In thick-walled phase, zooecial walls distinctly laminate, laminae arching orally convex, continuous across zooecial boundary zone. Light-colored granular tissue commonly forms small, rounded masses in outer cortex.

Intermediate diaphragms occurring in zooecia subjacent to intrazoarial overgrowths.

Taxa included. - Based on observations of internal characters seen in the primary types and identified specimens from the type localities, the following species originally included in Ceriopora by Goldfuss are considered to be correctly assigned to other genera: Ditaxia anomalopora (Goldfuss), Heteropora cryptopora (Goldfuss) and Zonopora spiralis (Goldfuss). Based on observations of the internal characters seen in identified specimens from the type locality, C. globosa Michelin is considered to be correctly assigned to Reptonodicava. Based on the observation of internal characters seen in identified specimens, C. dichotoma Goldfuss is tentatively referred to as Grammascoecia. The internal characters of other species assigned by others to Ceriopora are unknown to me.

Discussion. - Goldfuss (1826, p. 32) erected the genus Ceriopora and assigned 28 new species to it. He did not designate or indicate a type species. Gregory (1896, p. 195) listed the type species as $C$. micropora Goldfuss without comment, but later (Gregory, 1909, pp. 156-7) he indicated that his designation of lectotype followed, in part, the previous restrictions of Blainville (1830) and D'Orbigny (1854).

The concept of Ceriopora was based largely on two external characters (see Gregory, 1909, pp. 156-158 for an excellent review of development of the generic concept in the last century):

1) Aperture size - Blainville (1830, p. 378) separated Heteropora from Ceriopora. He stated that two kinds of zooecia could be recognized in Heteropora. Each polymorph was characterized by
the size of its aperture. Thus, Blainville implied that only one kind of zooecium could be recognized in Ceriopora. This generic distinction was followed by D'Orbigny (1854, p. 1029), Gregory (1896, p. 195; 1909, p. 156), Canu and Bassler (1920, p. 678), and Bassler (1953, p. G67). The concept of one kind of zooecium based on the size of the aperture was referred to as "monomorphic", or "lacking mesopores" by authors after D'Orbigny. The continued use of this character as generically diagnostic does not appear to be justified, however, because the frequency distributions of cross-sectional dimensions of zooecial chambers made from the type specimens of both Ceriopora micropora Goldfuss and Heteropora cryptopora (Goldfuss), their respective type species, are unimodal, and approach a normal distribution (Textfigs. 7A and 14A).
2) Intrazoarial overgrowth - species with overgrowths have been referred to as "multilamellar" (Canu and Bassler, and Gregory), or "plusiers couches superposées" (d'Orbigny). Overgrowths can often, but not invariably, be recognized on the external surfaces of zoaria, especially when the surface is slightly worn. De Blainville (1830) and D'Orbigny (1854) believed that zoaria of Ceriopora were composed of several overgrowths. D'Orbigny's earlier position was followed by Canu and Bassler (1926, p. 19) and Gregory (1909, p. 157). Gregory noted that "a certain amount of marginal lamellation must be expected in massive Bryozoa". Ceriopora, as based on observations of the type specimens, does have intrazoarial overgrowths and would thus be multilamellar in the sense of Blainville and D'Orbigny (1854).

The mode and amount of intrazoarial overgrowth is apparently a useful taxonomic character. The mode of growth shown in the type species of Ceriopora is intermediate in appearance between Reptonodicava and Heteropora. The type species of all three genera show a reduced endozone, apparently restricted to the proximal-most portion of encrusting zooecia, budding from a basal layer. All three are erect, massive, and globular to subramose in appearance. In Heteropora, intrazoarial overgrowth plays an important role in zoarial increase. The zoarium is composed of nested intrazoarial overgrowths connected by a few zooecia with continuous zooecial chambers, but
the zooecial walls from subjacent to superjacent growth commonly show some evidence of discontinuity of growth, such as great reduction in size and separation by a dark line.

In the type species of Reptonodicava overgrowths are rare, covering only a few zooecia at most, and zooecia commonly grew continuously in all directions for very long distances.

In the type specimen of Ceriopora intrazoarial overgrowth plays an intermediate role relative to Heteropora and Reptonodicava.

Gregory (1896, p. 95) included the statement, "Diaphragms horizontal, numerous", in his diagnosis of Ceriopora. This inclusion was based on his study of thin sections of Jurassic specimens which he assigned to Ceriopora globosa Michelin. The internal characters of C. globosa Michelin are sufficiently different from those of $C$. micropora to separate both in different genera. Bassler (1935, p. 186) designated $G$. globosa Michelin as the type species of Reptonodicava, but indicated that Reptonodicava was a synonym of Ceriopora. $R$. globosa does have numerous, closely spaced basal diaphragms, and the reference to numerous diaphragms in the definition of Ceriopora was reiterated by Canu and Bassler (1920, p. 678) and Bassler (1953, p. G67). Diaphragms occur in the lectotype of $C$. micropora, but they are intermediate diaphragms and no more than one was observed to occur within a single zooecium, unlike the numerous basal diaphragms typically seen in R. globosa.

Remarks on wall structure. - Light-colored, optically nearly structureless tissue in the zooecial wall is interpreted as being originally granular. The cortex of the zooecial walls in the endozone and thin-walled exozone portions are composed almost completely of granular tissue. Lamination becomes increasingly distinct in the thick-walled portions, and light-colored tissue is restricted to discontinuous bodies in the outer cortex alternating with, and surrounded by, laminated tissue. Masses of light-colored tissue bounded conformably by laminate tissue probably represent originally granular tissue ( Pl . 9, fig. 1b - lower, thin-walled portion of zooecial wall in center - 1c). Masses of light-colored tissue which cut across laminae are inferred to be recrystallized from originally (at least in part) laminate tissue ( Pl .9 , fig. 1b - upper, thick-walled portion of zooecial wall in center).

The wall structure is similar to that seen in Coscinoecia radiata Canu and Lecointre (Pl. 14, figs. 1e, f; Pl. 15, fig. 1g).

Ceriopora micropora Goldfuss, 1826 Pl. 7, figs. 1a-f; Pl. 8, figs. 1a-d; Pl. 9, figs. 1a-d
1826. Pars Ceriopora micropora Goldfuss, Petrefacta Germaniae, vol. 1, p. 33, pl. 10, figs. 4a, d; not figs. 4b, c.
1830. Ceriopora micropora Goldfuss, Blainville, Zoophytes: Dictionnaire de Science Naturelles, vol. 60, p. 378.
1834. Ceriopora micropora Goldfuss, Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 413.
1851. Pars Ceriopora micropora Goldfuss, von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, p. 52, pl. 5, fig. 13.
1909. Pars Ceriopora micropora Goldfuss, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 158-161.
1953. Pars Pennipora beyrichi Hamm, 1881, Voigt, Geol. Staatsinst. Hamburg, Mitt., Bd. 22, pp. 58, 61, 62, pl. 2, fig. 4.

Type. - UB 119 is designated as the lectotype. This specimen was figured by Goldfuss (1826, pl. 10, figs. 4a, d), von Hagenow (1851, pl. 5, fig. 13 as Heteropora crassa von Hagenow, 1851), Voigt (1953, pl. 2, fig. 13 as Pennipora beyrichi Hamm, 1881), and here (Pl. 7, figs. 1a-f; Pl. 8, figs. 1a-d and Pl. 9, figs. 1a-d).

Type locality and horizon. - "St. Petersberge bei Maastricht [Limbourg, Netherlands], aus dem Mergel bei Essen an der Ruhr [Federal Republic of Germany], und aus der Conchilienbreccie in der obern Schicht der Kreide von Cleom bei Nantu". Locality data were not listed with the figures, and Von Hagenow (1851) apparently did not find any locality data with the specimens. Commonly, Ceriopora micropora is listed as a Maastrichtian fossil from Maastricht, Netherlands. Many specimens from the area of Maastricht, externally similar to the lectotype, were sectioned. On the basis of internal characters, these specimens were assigned to Heteropora cryptopora or to two other species referable to other genera perhaps unnamed at present. Thus, reference of C. micropora to Maastrichtian at Maastricht, Netherlands, should be considered questionable.

Material studied. - The lectotype UB 119 was borrowed from Geol.-Paleont. Inst. Bonn University, West Germany. Three thin sections and three acetate peel replicas on a single acetate slide were prepared. Much of the specimen remained after sectioning. Dupli-
cate acetate peels are preserved in the National Museum of Natural History and the author's collection.

Description of the lectotype. - This description is based solely on the lectotype, the only specimen of $C$. micropora known to be available for study. This description includes assessment of nongenetic variation within this colony. No assessment of genetic or other interzoarial variation within C. micropora is implied in this description.

Endozone - The zooecial walls are undulatory to straight and thin (about .007 and .012 mm ). The walls have a granular cortex and a thin, dark zooecial lining (Pl. 7, fig. 1d; Pl. 8, figs. 1c, d).

Exozone - In the thin-walled exozone phase, zooecial walls are commonly thickened symmetrically across the boundary zone, and are submoniliform in profile. Monili are generally clavate to fusiform. Interzooidal pores are rare. Zooecial walls are homogeneous to slightly granular with a thin zooecial lining.

In the thick-walled exozone phases, zooecial walls are thickened symmetrically to asymmetrically across the boundary zone, and have moniliform cross sections. Successive monili are commonly unequal in length and thickness ( Pl . 9, figs. 1b, c), and sometimes show abrupt changes in growth direction among the monili (Pl. 9, fig. 1b). Zooecial chambers are commonly subelliptical in cross section, but show great variability (Table 7). Interzooidal pores are numerous (Text-fig. 7D).

Diaphragms - Intermediate diaphragms are occasionally observed, up to one per zooecium. The diaphragms commonly occur about . $1-.2 \mathrm{~mm}$ aboral to zooecial apertures, and subjacent to intrazoarial overgrowths (PI. 9, figs. 1a, b). The oral surfaces of the diaphragms are generally planar; the aboral surfaces are planar to strongly convex orally, and sometimes have an aborally flexed abutment which tapers into a short zooecial lining (Pl. 9, figs. 1a, b). Diaphragms are highly variable in thickness (Table 6, IntD-Th).


Text-figure 7 A-D. Histograms and cumulative curve from the lectotype of Ceriopora micropora Goldfuss. A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.
STATISTICAL SUMMARY OF MEASUREMENTS OF CERIOPORA MICROPORA GOLDFUSS

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zoarial |  |  |  |  |  |  |  |  |  |
| $\mathrm{Zr}-\mathrm{Ht}$ | 20 |  |  |  |  | 1 |  | 1 | 1 |
| Zr-Wth | 10 |  |  |  |  | 1 |  | 1 | 1 |
| Zooecial - Exozone |  |  |  |  |  |  |  |  |  |
| Zch-CsSn-NMxDn | . 16 | . 05 | . 11 | . 02 | 23 | 100 | 100 | 1 | 1 |
| ZcCh-CsSn-MxDn | 2.2 | 1.0 | 1.3 | 0.2 | 17 | 100 | 100 | 1 | 1 |
| Zche-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| $\mathrm{CdZcWl}-\mathrm{Th}$ | . 107 | . 008 | . 042 | . 018 | 42 | 100 | 100 | 1 | 1 |
| ZdPr-Cn/ZcCsSn | 4 | 0 |  |  |  | 100 | 100 | 1 | 1 |
| ZdPr-MnDr | . 014 | . 003 | . 009 | . 004 | 44 | 13 | 11 | 1 | 1 |
| Diaphragm-ExozoneIntD-Th $\quad .030 \quad .008 \quad .016 \quad .010 \quad 62 \quad 5$ |  |  |  |  |  |  |  |  |  |
| * In millimeters |  |  |  |  |  |  |  |  |  |
| 1. Lectotype UB 119 |  | EY To | ECIME | ODE |  |  |  |  |  |

TABLE 7
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN CERIOPORA MICROPORA GOLDFUSS

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Exozone <br> Regular |  | 17 | 7 |  |  |  |  |
| Sub | 2 | 38 | 8 |  | 3 |  |  |
| Irregular | 3 | 10 |  |  |  | 2 |  |
| Endozone <br> Regular |  | 4 | 4 |  |  |  |  |
| Sub | 1 | 10 | 4 | 1 |  |  |  |
| Irregular |  |  |  |  | 1 |  |  |

Discussion. - Designation of UB 119 as the lectotype agrees with restrictions first made by Blainville (1830, p. 378) and, in part, with restrictions in reference to figure 4 d , but not figure 4 a made by Von Hagenow (1851, p. 52) and followed by Gregory (1909, pp. 159-60) and Voigt (1953, pp. 58-62).

Von Hagenow (1851, pp. 48, 51) stated that he had studied three specimens which he believed were the original specimens before Goldfuss when he described Ceriopora micropora. The three specimens were attached to a small tablet ("Täfelchen") found in the Bonn Museum. Two were syntypic, as they were clearly identifiable with the specimens figured by Goldfuss; the first as plate 10 , figure $4 \mathrm{a}=\mathrm{UB} 119$, and the second as plate 10 , figures 4 b and c . Von Hagenow was of the opinion that neither of these two specimens was consistent with the concept of $C$. micropora. He was able to demonstrate that the specimen figured by Goldfuss as plate 10 , figures 4 b and c was a sponge, and he assigned UB 119 to his new species Heteropora crassa.

Von Hagenow believed that the third specimen on the small tablet and Goldfuss' plate 10, figure 4d were consistent with the concept of $C$. micropora and referred to both the specimen and the figure as the name-bearer for C. micropora. Furthermore, Von Hagenow believed that Goldfuss had taken the magnified view (pl. 10 , fig. 4d) from the third specimen; but as Gregory (1909, p. 160) noted: "Von Hagenow admitted an element of doubt in reference to fig. 4d."

Gregory (1909, pp. 159-60) and Voigt (1953, pp. 61-62) accepted Von Hagenow's restrictions and designation. Voigt (1953, pp. 58-64), however, believed that UB 119 is not conspecific with H. crassa Von Hagenow but is assignable to Pennipora beyrichi Hamm, 1881. These contentions are not here evaluated because of the present lack of knowledge concerning the internal characters of the respective types. Consideration of characters revealed in thinsections of specimens hitherto considered to be assignable to named species of such genera as Ceriocava, Ceriopora, Heteropora, Reptomulticava, and Reptonodicava, suggests that external characters are generally not diagnostic at the generic and family, and often not even at the species level.

For reasons listed below, Von Hagenow's contention that figure 4 d of Goldfuss was taken from the specimen figured by Von Hagenow (1851, pl. 5, fig. 4) is rejected.

1) Goldfuss clearly indicated that plate 10 , figures 4 a and 4 d were made from the same specimen by connecting them with a dashed line. Goldfuss and other authors commonly used the convention of connecting different views of the same specimen by a dashed line. Von Hagenow demonstrated that figures $4 b$ and 4 c of Goldfuss' plate 10 were different views of the same specimen; both views were connected by a dashed line. Figures 4a and 4d should be considered as different views of UB 119 unless strong evidence to the contrary were to be offered.
2) The lectotype is close enough in appearance for acceptable comparison with the magnified and somewhat idealized view of the surface as figured by Goldfuss (pl. 10, fig. 4d).
3) Von Hagenow was not positive of equating the Goldfuss figure (pl. 10, fig. 4d) with the third specimen on the card (Von Hagenow, pl. 4, fig. 5), and Von Hagenow emphasized that only the figure by Goldfuss and the third specimen fitted the concept.
As outlined above, Von Hagenow is believed to have made a mistake in equating the specimen he figured as plate 5 , figure 4 , with plate 10 , figure 4 d of Goldfuss, and thus he designated two specimens to be the name bearer. Von Hagenow's elimination of the sponge specimen (Goldfuss, pl. 10, figs. 4c, d) from consideration is here followed, and the lectotype is designated as the remaining figured specimen, UB 119 figured by Goldfuss (pl. 10, figs. 4 a and 4d).

Remarks on enigmatic structures. - A single, unusual dia-phragm-like structure was observed in the transverse section and is illustrated in Pl. 9, figure 1d. This structure, known only from a single profile view, apparently forms a hollow, tubelike structure within the zooecial chamber which flares, and is attached by its oral and aboral extremities to the zooecial wall. The structure is considered to be primary because of its structural continuity with the zooecial wall.

Cross sections of two large chambers were observed in tangential section (one is figured in Pl. 8, fig. 1a). The cross-sectional dimensions are more than twice as long as the largest zooecia observed (Table 6), and are interpreted as interzooecial structures. The walls are unbroken and are interpreted as primary in origin. The only large primary chambers in other cerioporids are thought to be brood chambers. In lieu of better evidence, these inter-zooecial spaces in $C$. micropora are interpreted as brood chambers.

Genus CORYMBOPORA Michelin, 1846
Type species: Corymbopora menardi Michelin, 1846, p. 213, pl. 53, fig. 10 by monotypy.
1846. Corymbopora Michelin, Iconographie Zoophytologique, description par localités et Terrains des Polypiers Fossiles de France et pays environnants, p. 213.
1850. Pars Fasciculipora d'Orbigny (1846), D'Orbigny, Prodrome de Paléontologie stratigraphique universelle, vol. 2, p. 177.
1854. Corymbosa d'Orbigny, Terrain Crétacé Bryozoaires: Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 691.
1890. Pars Fasciculipora d'Orbigny (1846), Pergens, Soc. Belge Géol. Paleont. Hydro. Bull., vol. 3, p. 377.
1909. Corymbopora Michelin, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 45.
1916. Non Corymbopora Lang, Ann. Mag. Nat. Hist., ser. 8, vol. 18, p. 382.
1919. Corymbopora Michelin, Canu, Soc. Géol. France, Bull., vol. 17, ser. 4, p. 348.
1953. Corymbopora Michelin, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G70.

Tentative diagnosis. - Zoaria branched. Zooecia dimorphic. Large zooecia relatively long, growing parallel to branch axis, and opening only at distal growing tips of branches. Small polymorphs short, budding obliquely from walls of most laterally-disposed zooecia in branch. Apertures of small polymorphs cover side of branch.

Zooecial walls of large polymorphs granular to indistinctly laminate.

Zooecial walls of small dimorphs indistinctly laminate. In zooecial wall, between longitudinally adjacent small polymorphs, laminae form saddle-shaped configurations which arch orally convex longitudinally, but sag transversely, forming aborally convex arches. In zooecial wall, between laterally adjacent small polymorphs, laminae diverge orally at low angles from zooecial boundary zone.

Both intermediate and basal diaphragms occur.
Taxa included. - The internal characters of $C$. neocomiensis d'Orbigny, based on an examination of topotype specimens, are consistent with the concept of Corymbopora.

Fungella, as based on an examination of identified specimens of the type species $F$. dujardini von Hagenow, is closely similar to Corymbopora in growth habit and, perhaps, other characters. Assessment of these apparent similarities must await a more detailed examination of types, or at least topotypes, of $F$. dujardini. Internal characters of other species assigned to Corymbopora are unknown to me.

Discussion. - D'Orbigny (1854, p. 689) recognized dimorphism in Corymbopora: "Chaque branch est terminée par un gros faisceau de cellules verticales . . . La paroi externe des faisceaux est partout criblée en long, de nombreux pores intermédiaires par lignes longitudinales". Gregory (1909, p. 44) noted that there were two sizes of zooecia, but stated that "the sides of the stem are covered by an epizoarial layer, marked by numerous pores, the remnants of the aperture of dead zooecia".

D'Orbigny (1850) and Pergens (1890) synonymized Corymbopora with Fasciculipora d'Orbigny (1846). D'Orbigny (1854) recognized Corymbopora and Fasciculipora but assigned both to the family Fasciporidae d'Orbigny. Gregory (1909) included Corymbopora in the Fascigeridae d'Orbigny in which he also included Fasciculipora. Borg (1926a) included the genera Domopora and Defrancia in the family Corymboporidae Smitt. Furthermore, Borg (1926a) assigned the Corymboporidae and Fasciculipora to his division 2, the Acamptostega (approximately equivalent to the suborder Tubuloporina Milne-Edwards, 1838), which are characterized, in part, by single-walled growth of zooecial walls and pseudopores.

In all of the above genera, growth habit is similar in that large zooecia grow nearly parallel to the growth axis of the branch throughout their length, and apertures are located only at the distal ends of branches. On the other hand, pseudopores, not small dimorphs, are reported in Fasciculipora (Borg, 1926a, p. 19). The internal characters of Domopora and Defrancia are practically unknown, but illustration of external features of these genera (Borg, 1926a, text-fig. 49, p. 300; text-figs. 79-80, p. 378; text-figs. 83-85, p. 383) do not show the presence of small dimorphs characteristic of C. menardi Michelin, nor were they reported.

The wall structure observed in Corymbopora is consistent with Borg's double-walled concept, but Corymbopora, as presently understood, is not easily assignable to Borg's double-walled groups, the Pachystega ( $=$ Horneridae), or the Calyptrostega ( $=$ Lichenoporidae). The author has followed Bassler (1953) in discussing Corymbopora with other genera assigned to the Cerioporina ( $=$ Heteroporina of Borg), the third group consistent with Borg's double-walled hypothesis. Thin-sections of specimens assigned to Fungella and Corymbopora neocomiensis Canu and Bassler reveal small dimorphs covering the stem, and other structures suggesting affinity with $C$. menardi. Detailed study of these species and others is needed before a formal taxonomic action is made, such as possible reassignment of species and erection of a new group equivalent to Borg's other double-walled divisions.

Corymbopora menardi Michelin. 1846 Pl. 10, figs. 1a-c, 2, 3a-d, 4a-b;
Pl. 11, figs. 1a-d, 2, 3a-b; Pl. 12, figs. 1, 2a-d
1846. Corymbopora menardi Michelin, Iconographie Zoophytologique, description par localités et Terrains des Polypiers Fossiles de France et pays environnants, p. 213, pl. 53, fig. 10.
1854. Corymbosa menardi (Michelin), d'Orbigny, Terrain Crétacé Bryozoaires: Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 691, pl. 744, figs. 7-12. Obj. syn.
1909. Corymbopora menardi Michelin, Gregory, Catalogue of fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 45.

Type. - A holotype was not designated by Michelin, nor has a lectotype been designated since. Michelin's primary types are thought to have been originally placed in the Caen Museum, or possibly the Muséum National d'Histoire Naturelle. The museum at

Caen was destroyed in the Second World War, and none of Michelin's specimens of Corymbopora menardi have been found in the collections of the Muséum National d'Histoire Naturelle (pers. comm. E. Buge, 1969).

Type locality and horizon. - Le Mans (Sarthe), France, Cretaceous, Cenomanien.

Material studied. - Prof. E. Buge kindly made available specimens collected by Ferdinand Canu from the topotype locality. The suite of specimens contains a single, nearly complete zoarium, MNHN Canu Coll. 57057-1; and a number of zoarial fragments, MNHN Canu Coll. 57057-2. Thin sections and acetate peels were made of six specimens from MNHN Canu Coll. 57057-2. The original labels indicate that the specimens were collected from "Cenomanien, Le Mans" (Sarthe), France, the type locality. The specimens, MNHN Canu Coll. 57057-2, are labeled as "Fasciculipora (Corymbosa) menardi d'Orb." In addition, thin sections and acetate peels were prepared from four specimens in the United States National Museum collection. The specimens, USNM Loc. 2947, were collected from the type locality. Duplicate acetate peels of all specimens sectioned are preserved in the United States National Museum collection and the author's collection.

Description. -
Mode of growth — Zoaria are small (see Table 8, Zr-Ht, ZrWth); branching is dichotomous, producing a delicate, arborescent architecture. Branches have circular cross sections, but expand distally to form capitate distal tips (Pl. 7, fig. 1b) with circular to elliptical cross sections (Pl. 7, fig. 1a).

Endozone - Zooecia have thin, parallel-sided walls which are locally undulatory but, in general, grow parallel to the growth axis. Walls are homogeneous to subgranular and have thin, dark-colored zooecial linings.

Exozone - Remnant, thick-walled growth phases are seen in the capitular growing tips. They are commonly located just proximal or distal to a brood chamber (Pl. 10, fig. 3b). The zooecial walls are annularly thickened. The thickenings are roughly symmetrical across adjacent zooecia forming a zone parallel to the zoarial surface ( Pl. 10, fig. 3b). Longitudinally, the zooecial walls have monili-
form profiles, symmetrical in thickness across the boundary zone. Thick-walled zones are thin, and commonly have one to four monilar thickenings longitudinally.

Small polymorphs - Apertures of small polymorphs are arrayed in longitudinal rows on the branch surface. The portion of the zooecial walls between transversely adjacent zooecia forms prominent ridges parallel to the growth axis of the branch (Pl. 10, figs. 1b, 2). Small polymorphs have uniformly short, reflexed, conical, or sockshaped chambers ( Pl. 11, figs. 1b, c, e). Interzooidal pores pass through the proximal wall to large polymorphs (Pl. 11, fig. 1e), and also through the walls of longitudinally adjacent zooecia (Pl. 11, fig. 1b).

Diaphragms - Basal diaphragms were occasionally observed in endozones of large polymorphs. The diaphragms are relatively planar, and show slight oral flexure at the intersection with the zooecial wall. Diaphragms commonly were observed at approximately the same level in several adjacent zooecia.

Intermediate diaphragms were observed rarely. The diaphragms flex aborally and merge with the zooecial lining.

Brood chambers - Brood chambers are abundant (up to ten were observed in a single zoarium). Abandoned brood chambers were observed only in capitular areas, and are associated with thickwalled zones. Brood chambers are lobate; several lobes branch from a single main lobe, continuous proximally with a single zooecial aperture (Pl. 12, fig. 1). Subjacent zooecia are sealed by a thin floor which is structurally continuous with subjacent zooecial walls ( Pl . 12, figs. 2d, e). Brood chambers observed at the zoarial surface were not roofed over. Abandoned brood chambers were covered by a distal wall bearing interzooidal pores, and growing continuously from the lateral compound wall (Pl. 12, figs. 2d, e.) Zooecia budding from the distal wall are commonly thick-walled and have moniliform profiles (Pl. 12, figs. 2d, e).

Discussion. - Athough Michelin's primary types were not available for examination, specimens were assigned to $C$. menardi with confidence because:

1) All specimens can be recognized on the basis of external characters, zoarial shape, distally expanded or capitate branches, and
the arrangement of the apertures of small polymorphs. These characters are well shown in Michelin's figure.
2) The species concept, as developed in descriptions or illustrations of D'Orbigny (1854), Gregory (1909), and Canu (1919) has been consistent.
3) Specimens studied by D'Orbigny (1854), Gregory (1909), Canu (1919), and herein were collected from the type locality.

Under these circumstances, there are no taxonomic difficulties imposed by the unavailability of the type specimens. Thus, by Art. 7 (a, i), p. 81, ICZN, 1964, there is no necessity for designating a neotype.

Remarks on brood chambers. - In Corymbopora menardi Michelin there is some indication that the brood chamber maintained a connection with a single, presumably fertile, zooecium and hence might be termed a gonozooecium. The use of this term, however, implies that the whole structure is a zooecial homologue. This is difficult to apply because the wall which bounds the brood chamber is compound, and shared with subjacent and lateral zooecia. The depositing epithelium is probably better considered as a zoarial rather than a zooecial tissue. For this reason, the more general term, brood chamber, is here retained.

In $C$. menardi, the brood chamber was never roofed over by a porous, diaphragm-like structure typical of the brood chambers in Ceriocava corymbosa (Lamouroux), Heteropora cryptopora (Goldfuss) and Parleiosoecia jacksonica Canu and Bassler. The brood chambers in C. menardi were apparently covered by calcareous tissue only after abandonment. The covering structure is similar in appearance to the adjacent zooecial wall rather than the basal, layerlike structure seen in other genera.

Genus COSCINOECIA Canu and Lecointre, 1934
Type species: Coscinoecia radiata Canu and Lecointre, 1934, p. 21, by original designation and monotypy.
1934. Coscinoccia Canu, and Lecointre, Les Bryozoaires Cyclostomes Des Faluns de Touraine et d’Anjou, Soc. Géol. France, Mém., vol. 9, No. 4, p. 198.
1953. Coscinoccia Canu and Lecointre, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G70.
1957. Coscinoecia Canu and Lecointre, Buge, Mus. Nat. d'Hist. Nat., Mém., ser C, Sciences de la Terre, vol. 6, p. 121.
TABLE 8
sTATISTICAL SUMMARY OF MEASUREMENTS OF CORYMBOPORA MENARDI MICHELIN

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} \mathrm{rr}^{\text {r }}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zoarial |  |  |  |  |  |  |  |  |  |
| Zr - Ht | 13. |  |  |  |  |  |  |  | 6 |
| Zr-Wth | 11. |  |  |  |  |  |  |  | 6 |
| Zooecial-Large Polymorphs - Endozone |  |  |  |  |  |  |  |  |  |
| ZcCh-CsSn-MxDn | . 20 | . 08 | . 14 | . 02 | 17 | 125 | 125 | 3 | 1 |
| Zch-CsSn-NMxDn | . 17 | . 04 | . 11 | . 02 | 19 | 125 | 125 | 3 | 1 |
| ZcCh-CsSn-MxDn | 2.1 | 1.0 | 1.3 | . 22 | 17 | 125 | 125 | 3 | 1 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| $\mathrm{CdZcWl}-\mathrm{Th}$ | . 022 | . 005 | . 010 | . 003 | 31 | 125 | 125 | 3 | 1 |
| ZdPr-Cn/ZcCsSn | 5. | 0. |  |  |  | 125 | 125 | 3 | 1 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF CORYMBOPORA MENARDI MICHELIN

| Character |  |  | X* | S* | C. V. | N | NZc | $\mathrm{N} \mathbf{Z r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ```Zooecial - Small Polymorphs ZcCh-CsSn-LngDn``` | . 07 | . 04 | . 05 | . 01 | 19 | 12 | 12 | 2 | 2 |
| Zch-CsSn-TrvDn | . 06 | . 04 | . 05 | . 01 | 12 | 12 | 12 | 2 | 2 |
| CdZcWl-ThLng | . 05 | . 02 | . 04 | . 01 | 21 | 12 | 12 | 2 | 2 |
| CdZcWl-ThTrv | . 07 | . 03 | . 05 | . 02 | 33 | 16 | 16 | 2 | 2 |
| Brood Chamber BrCh-WthLb | . 90 | . 45 | . 60 | . 03 | 7 | 18 |  | 2 | 4 |
| BrCh-Dth | 2.2 | . 8 | 1.5 | . 3 | 4 | 20 |  | 4 | 5 |
| *In millimeters |  |  |  |  |  |  |  |  |  |
| KEY TO SPECIMEN CODE |  |  |  |  |  |  |  |  |  |
| 1. Topotypes: MNHN Canu Coll. 57057-2 (3), 50; (5), 25 ; (7), 50. |  |  |  |  |  |  |  |  |  |
| 2. Topotypes: MNHN Canu Coll. 57057-2 (3), (6) ; (4) (6). |  |  |  |  |  |  |  |  |  |
| 3. Topotypes: MNHN Canu Coll. 57057-2 (3) (14) ; (4) (8) ; (5) (13). |  |  |  |  |  |  |  |  |  |
| 4. Topotypes: MNHN Canu Coll. 57057-2 (6), (8) ; (7), (10). |  |  |  |  |  |  |  |  |  |
| 5. Topotypes: MNHN Canu Coll. 57057-2 (1) (4); (4), (7) ; (5), (2); (6), (7). |  |  |  |  |  |  |  |  |  |
| 6. Topotype: MNHN Canu Co |  |  |  |  |  |  |  |  |  |

Tentative diagnosis. - Zoaria branched; branches have welldifferentiated coaxial endozones and exozones. Zooecia dimorphic. Large polymorphs occur in intermonticular areas, small polymorphs in intermonticular and monticular areas.

In exozone, zooecial walls distinctly laminated. Laminae broadly arched across zooecial boundary zone. Laminate tissue sometimes alternates with granular tissue which forms discontinuous rounded masses in outer cortex.

Basal diaphragms numerous and closely spaced in endozone and zone of flexure. Intermediate diaphragms occur in intrazoarial overgrowths.

Taxa included. - Monotypic for the type species, $C$. radiata.
Discussion. - Coscinoecia is a large bryozoan, distinctive both externally and internally. Bassler (1953, p. G70) assigned Coscinoecia to the family Tetrocycloeciidae Canu and noted in the diagnosis that Coscinoecia is "like Tretocycloecia [sic] but oeciostome larger than zooecial apertures". Coscinoecia is similar to Tetrocycloecia, Leiosoecia, and Ceriocava in its coaxially ramose growth habit. Coscinoecia, however, differs from all three genera in several characteristics, such as the structure and appearance of the zooecial walls, the arrangement of dimorphs, and the occurrence of diaphragms. The gross morphologic structures are consistent with placement in the Cerioporina, but the differences in morphology are too great, as presently understood, to indicate close affinities with any genus studied here.

Coscinoecia radiata Canu and Lecointre, 1934
Pl. 13, figs. 1a-h;
Pl. 14, figs. 1a-h; Pl. 15, figs. 1a-i
1934. Coscinoccia radiata Canu, and Lecointre, Soc. Géol. France, Mém., vol. 9 No. 4, p. 198, pl. 40, figs. 1-7.
1957. Coscinoecia radiata Canu and Lecointre, Buge, Mus. Nat. d'Hist. Nat., Mém., ser. C, Sciences de la Terre, vol. 6, p. 122, No fig.
Type. - Canu and Lecointre (1934) noted that they had examined several specimens from seven localities in the Helvetien of southern France. One specimen from the primary type suite has been located. This specimen, MNHN Canu Coll. 58872, is the large, well-preserved specimen figured by Canu and Lecointre (1934, pl. 40, figs. 1-4) and is here designated as the lectotype. Thin-sections were figured by Canu and Lecointre (pl. 40, figs. 5-7); the specimen
or specimens sectioned were not identified. The thin-sections were not located by the author. Although Buge (1957, p. 122) referred to MNHN Canu Coll. 58872 as the holotype, there is no such designation in the primary source.

Type locality and horizon. - Canu and Lecointre (1934, p. 198) cited the following localities in France for the occurrence of $C$. radiata:

Faluns (Helvetien)
Indre-et-Loire: Saint Epain, Ferrière-l'Arcon
Marn-et-Loire: le Haguineau, Doué-la-Fontaine, Montlouet
Ille-et-Vilaine: environs de Rennes
Burdigalien supérieur
Piémont: Croce Barton
The label with the lectotype lists the word Doué; therefore, the type locality and horizon are restricted to the Faluns (Helvetien) [Miocene], Doué-la-Fontaine, Marn-et-Loire [France].

Material studied. - The lectotype was kindly loaned to the author by Dr. E. Buge, Muséum National d'Histoire Naturelle. Five thin-sections and 13 acetate peel replicas were made from the lectotype; most of the original zoarial fragment and three remnants remain after sectioning. Duplicate peels are in the United States National Museum and the author's collection.

Description of the lectotype. - This description is based solely on the lectotype, the only specimen of $C$. radiata known to be available for study. This description includes assessment of some nongenetic variation within this colony. No assessment of genetic or other interzoarial variation within C. radiata is implied in this description.

Mode of grozuth. - The zoarium is robust (Table 9, $\mathrm{Zr}-\mathrm{Ht}, \mathrm{Zr}-$ Wth, Br-CsSn-MxDn); branches are roughly cylindrical. Most of the surface of the primary branch is encrusted by an intrazoarial overgrowth (Pl. 13, figs. 1e, f).

Large polymorphs occur in intermonticular areas, and are arranged in rows disposed radially about the monticular areas ( Pl . 13, figs. 1b, c). Small polymorphs occur in intermonticular areas, commonly in rows between the large polymorphs (Pl. 13, figs. 1b, c), and also in monticular areas ( Pl .13 , figs. 1b, c, h; Pl. 14, fig. 1d). Less regular arrangements occur locally (Pl. 13, fig. 1h).

Endozone. - Zooecial walls are thin, symmetrically to subsymmetrically thickened across the boundary zone, and have submoniliform profiles. Zooecial chambers commonly have subelliptical, subpolygonal cross sections. Zooecial walls are generally granular, sometimes indistinctly laminate, and a thin zooecial lining is usually present. Basal diaphragms are commonly thin and planar to slightly convex aborally. At the juncture with the zoecial wall, diaphragms flex orally and merge with the zooecial lining ( Pl .13 , fig. 1d; Pl. 15 , figs. lb, i).

Exozone. - The zooecial walls of adjacent large polymorphs are symmetrically to subsymmetrically thickened across the zooecial boundary zone and have moniliform profiles, largely because of the occurrence of numerous, widely-flexed, interzooidal pores (Pl. 13, fig. 1 g ). Monilar profiles commonly are elliptical, ovate or fusiform, but show less variation in thickness longitudinally than small polymorphs, and appear nearly parallel-sided ( Pl . 13, figs. 1e, g). Zooecial chambers commonly have subelliptical cross sections but show relatively large variation in this respect (Table 10). Cross sections are often somewhat irregular because of crescentic inflections in the zooecial wall (Pl. 14, fig. 1h).

Zooecial walls of small polymorphs are commonly thickened annularly. Back-to-back adjacent small polymorphs are thickened symmetrically across zooecial boundary zones (Pl. 13, figs. 1e, g; Pl. 14, figs. 1a, b, e), but are commonly thickened asymmetrically across zooecial boundary zones when adjacent to large polymorphs ( Pl . 13, fig. 1g; Pl. 14, figs. 1b, f; Pl. 15, fig. 1g). Monilar profiles of adjacent small polymorphs are circular, elliptical, clavate, inversepyriform or alate. In monticular areas, cross sections of small polymorph chambers are subelliptical to subcircular (Pl. 14, fig. 1d). In intermonticular areas, cross sections of small polymorph chambers are commonly more elliptical and less regular. Small, blunt mural spines were often observed in both large and small polymorphs ( Pl . 14, figs. 1d, g).

Diaphragms. - Basal diaphragms were observed only in the endozone and zone of zooecial bending. The diaphragms are numerous, closely spaced (Table 9, Bsl-Intvl), and generally thin (Table 9, BsID-Th). The diaphragms are planar to slightly convex
TABLE 9
STATISTICAL SUMMARY OF MEASUREMENTS OF COSCINOECIA RADIATA CANU AND LECOINTRE

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\underset{\text { Zr-Ht }}{\text { Zoarial }}$ | 50 |  |  |  |  | 1 |  | 1 | 1 |
| Zr-Wth | 40 |  |  |  |  | 1 |  | 1 | 1 |
| Br-CsSn-MxDn | 11 | 5 |  |  |  | 2 |  | 1 | 1 |
| Zooecial-All Polymorphs - Exozone $\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}$ | . 24 | . 05 | . 12 | . 04 | 32 | 100 | 100 | 1 | 1 |
| Zch-CsSn-NMxDn | . 17 | . 03 | . 10 | . 04 | 37 | 100 | 100 | 1 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{Csin}-\mathrm{NM} M \mathrm{Dn}}$ | 3.3 | 1.0 | 1.3 | . 36 | 27 | 100 | 100 | 1 | 1 |
| CdZcWl-Th | . 160 | . 010 | . 052 | . 029 | 55 | 100 | 100 | 1 | 1 |
| ZdPr-Cn/ZcCssn | 3 | 0 |  |  |  | 100 | 100 | 1 | 1 |

*In millimeters
Statistical summary of measurements of Coscinoecta radiata canu and lecointre

| Character | O.R.* |  | $\mathrm{X}^{*}$ | S* | C. V. | N | NZc | $\mathrm{N} \mathrm{Zr}_{r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zooccial - All Polymorphs - Exozone ZcSp-Cn/ZcCsSn | (on.) | 0 |  |  |  | 100 | 100 | 1 | 1 |
| ```Zooecial-All Polymorphs - Endozone ZcCh-CsSn-MxDn``` | . 18 | . 04 | . 15 | . 04 | 24 | 25 | 25 | 1 | 1 |
| ZcCh-CsSn-NMxDn | . 16 | . 04 | . 12 | . 03 | 25 | 25 | 25 | 1 | 1 |
| $\frac{Z c \mathrm{Ch}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NM} \mathrm{M} \mathrm{Dn}}$ | 1.5 | 1.0 | 1.2 | . 13 | 11 | 25 | 25 | 1 | 1 |
| CdZcWl-Th | . 019 | . 005 | . 012 | . 004 | 35 | 25 | 25 | 1 | 1 |
| ZdPr-Cn/ZcCsSn | 2 | 0 |  |  |  | 25 | 25 | 1 | 1 |
| $\begin{aligned} & \text { Zooecial-Large Polymorphs } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 18 | . 08 | . 14 | . 02 | 16 | 57 | 57 | 1 | 1 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF COSCINOECIA RADIATA CANU AND LECOINTRE

STATISTICAL SUMMARY OF MEASUREMENTS OF COSCINOECIA RADIATA CANU AND LECOINTRE

aborally, and flex orally to form abutments before merging obscurely with the zooecial lining (Pl. 15, figs. 1b, i).

Most zooecia in the intrazoarial overgrowth have a single, thin (Table 9, IntD-Th) intermediate diaphragm at approximately .2 mm aboral to the skeletal aperture ( Pl .15 , fig. 1a). The diaphragms are generally planar, and they flex aborally to form a short abutment which adjoins, but does not merge with, the zooecial wall.

Brood chambers. - Brood chambers are large (Table 9, BrChLth, BrCh-Wth, BrCh-Dth) zoarial cavities with lensoid to rectangular profiles (Pl. 15, figs. 1c, f) and sublobate cross sections (Pl. 15 , fig. 1e). The floor is incomplete (Pl. 15, fig. 1c), and subjacent zooecia show various degrees of closure, from fully open to completely sealed. Closure may be made both by lateral growth of the apertural portion of the zooecial wall, and by emplacement of a nonperforate intermediate diaphragm ( Pl .15 , fig. 1h). The roof is apparently nonporous (Pl. 15, fig. 1c). Small polymorphs bud from a thin basal layer which encrusts the roof (Pl. 15, fig. 1d) forming monticular areas. The apertures of these small polymorphs are commonly elliptical to slitlike, in contrast to the more rounded openings of normal monticular areas. Encrusted brood chambers commonly open to the surface by means of a single, large, subcircular to elliptical opening (Pl. 13, figs. 1b, c).

TABLE 10
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN THE LECTOTYPE OF COSCINOECIA RADIATA CANU AND LECOINTRE

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| Exozone <br> Regular | 1 | 28 | 7 | 3 |  | 1 |  |
| Sub | 3 | 31 | 11 |  | 1 |  |  |
| Irregular | 3 | 4 | 2 |  |  |  | 5 |
| Endozone <br> Regular |  | 2 | 1 |  |  | 1 |  |
| Sub | 1 | 10 | 3 |  | 5 | 1 |  |
| Irregular |  |  |  |  |  |  | 1 |



Text-figure 8 A-D. Histograms and cumulative curve from the lectotype of Coscinoecia radiata Canu and Lecointre. A. Normal to maximum crosssectional dimension of a zooecial chamber. B. Ratio of the maximum crosssectional dimension of a zooecial chamber to the normal to maximum crosssectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.

Remarks concerning morphology. -
Growth habit - The erect, robust growth habit of C. radiata provided a suitable substrate for other bryozoan colonies. Several small cheilostome colonies and one lichenoporid colony were observed on the zoarial surface.

Intrazoarial overgrowth could be interpreted, in the situation figured in Plate 13, figures 1e, f , as a mechanism of defense because the overgrowth encrusts a relatively large cheilostome colony which had submerged part of the main stem.

Polymorphism - The starlike pattern engendered by the arrangement of dimorphic zooecia is a distinctive external character. In tangential sections the pattern is not always so readily visible and requires comment. Remarks on this subject must be considered tentative, however, because of limited material.

When viewing the zoarial surface, the eye readily makes the distinction between large and small polymorphs. Large polymorphs have relatively constant zooecial apertures in the exozone, and one sees a large, black zooecial opening ( Pl .13 , figs. 1b, c). On the other hand, small polymorphs have much more variable cross-sectional areas because of pronouonced annular thickening of the walls. When the zoarial surface is viewed, the eye "sees" the smallest cross section of small polymorphs as a black void; indeed, the largest dimension is nearly ignored unless an effort is made to see it (note light-colored areas around black voids at top, fig. 1c in Pl. 13).

In sections, zooecia are cut at all levels. If the section intersects small polymorphs where walls are thickest, the polymorphs are easily distinguishable. Conversely, where walls are thin, the areal dimensions approach, and perhaps overlap, the large polymorphs (see Text-fig. 8A). Thus, the regular pattern, so distinct at the zoarial surface, is obscure in thin-section.

Brood chambers - In comparison to brood chambers in Ceriocava, Heteropora and Parleiosoecia, Coscinoecia has brood chambers with nonporous roofs and incompletely calcified floors.

Another difference is that a single, large connection with the surface is retained well after the calcareous brood chamber roof was deposited, suggesting that larvae may have been contained, or brooded, after the chamber was roofed over and submerged beneath new growth.

## Genus DIPLOCAVA Canu and Bassler, 1926

Type species: Diplocava incondita Canu and Bassler, 1926, p. 71, by original designation.

## 1926. Diplocava Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 67, p. 71. 1953. Diplocava Canu and Bassler, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G71.

Tentative diagnosis. - Zoaria massive to branching. Growth habit variable, with complete gradation between two end members. In one growth habit, most zooecia relatively short and growing radially to produce lenticular intrazoarial overgrowths. In second habit, axial zooecia longer, initially growing nearly parallel to major growth axis before bending towards zoarial surface, and producing coaxial appearance; but, zooecial walls commonly not showing typical endozonal appearance in axial region.

In exozone, zooecial walls laminate. Laminae diverge orally from generally dark-colored, narrow boundary zone; then arch convex orally and recurve aborally. Cortex laminae abut thin zooecial lining at low angle. Walls commonly have integrate appearance in tangential section.

Simple external walls common. Intermediate and basal diaphragms occur.

Taxa included. - Only the type species was studied. Internal characters of other species assigned to Diplocava are unknown to me.

Discussion. - Diplocava was originally defined as "Ceriocavidae with dimorphic zooecia" (Canu and Bassler, 1926, p. 71). The dimorphs were described by Bassler (1953, p. G71) as " . . . large open ones separated by a zone of small ones with facets." Examination of thin-sections reveals, however, that zooecia are monomorphic. Another interpretation of the divergent zooecial appearance seen in Diplocava is given below.

The emplacement of simple external walls apparently represents a late phase in zooecial ontogeny. The portions of the compound zooecial wall adjacent to the external simple wall are consistent in having thick walls with recurved laminae, in having a well-defined boundary zone, and in having nearly parallel-sided profiles, characteristics not seen in more aboral portions of the zooecial wall.

In addition, zooecia of each type occur in groups. Zooecia with large apertures encrust the zooecia sealed by external walls ("small
ones with facets") (Pl. 16, fig. 1b). The present interpretation suggests that the "large zooecia" were in a stage of active, orally-directed growth, while the "small zooecia" had ceased active, orally-directed growth of the zooecial wall and had secreted simple external walls at the skeletal aperture.

The Ceriocavidae were defined by Canu and Bassler (1922, p. 89) on the appearance of the brood chamber which they described as "long, transverse, convex, regular, symmetrical vesicle with special walls". Although Canu and Bassler (1926, p. 71) noted in their remarks on the type species that the brood chamber was unknown in Diplocava, they described (on the same page) a "star-shaped" brood chamber from a "cotype" of the type species. This specimen is considered here to be assignable to another, perhaps unnamed, genus. Remarks concerning its morphology, i.e., brood chamber, are not applicable to the concept of Diplocava.

The assignment of Diplocava to the Ceriocavidae Canu and Bassler, based on the brood chamber alone, is not justified. Presently, lack of knowledge of changes in skeletal structure through time in cyclostomes makes phylogenetic inferences speculative; however, inferences of close relationship between Ceriocava and Diplocava cannot be supported by morphologic evidence as understood here. The respective type species differ in growth habit, wall structure, and occurrence of diaphragms. They differ, as well, in other characters tentatively considered significant at the specific level, e.g., profiles of zooecial walls, and outlines of zooecial chambers.

Simple external walls have been recognized in Haploecia. Diplocava differs from Haploecia in mode of growth and wall structure. Diplocava and Reptonodicava both have recurved laminae in the outer exozone. Reptonodicava differs from Diplocava in mode of growth and occurrence of diaphragms.

Diplocava incondita Canu and Bassler, 1926 Pl. 16, figs. 1a-h; Pl. 17, figs. 1-4, 5a-c, 6a, b; Pl. 18, figs. 1-3; Pl. 19, figs. 1-3
1926. Diplocava incondita Canu and Bassler, U.S. Nat. Mus., Proc., vol. 67, pp. 71-73, pl. 10, figs. 5 (specimen in lower right corner of figure), 6.
Type. - USNM 69925-2 figured by Canu and Bassler (1926, pl. 10, figs. 5 - specimen in lower right corner of photograph - and 6 ) is designated as the lectotype. Five unnumbered thin-sections labeled type are paralectotypes.

Type locality and horizon - Lower Cretaceous (Valangian), Sainte Croix (Vaud), Switzerland.

Material studied. - Five thin-sections and five acetate peels on one slide were made from the lectotype. In addition, 22 thin-sections and 24 acetate peels were made from 10 specimens collected from the type locality (USNM Loc. 2404) and identified as Diplocava incondita by Bassler. Duplicate acetate peels are preserved in the bryozoan collection of the National Museum of Natural History and in the author's collection.

Description. -
Mode of growth - Zoaria are small (maximum observed length is about 11 mm , diameter about 5 mm ), and are either irregularly domal to digitate masses or are branched with subcylindrical branches. The irregular zoarial appearance (Pl. 16, fig. 1a) is caused by the irregular repetition of both encrusting and coaxial modes, and by the variation in position and direction of the major growth axis from one overgrowth unit to another.

Intrazoarial overgrowths sometimes cover most of the surface of subjacent growth units (Pl. 17, fig. 3), or are restricted to the surface peripheral to the continuously-growing zooecia of the axial region (Pl. 16, fig. 1d; pl. 17, fig. 5a). Basal layers are generally thick (Pl. 16, fig. 1g; Pl. 17, figs. 5b, c, 6a, b). Proximal parts of overgrowths sometimes display thin, recumbent, zooecial walls interpreted as endozones (Pl. 16, fig. 1h, on left); but commonly, zooecial walls are essentially thick-walled throughout.

Exozone - The zooecial walls are symmetrically thickened across the boundary zone. The walls generally show a regular increase in thickness orally. Sometimes, however, the walls show a slight thinning in medial portions ( Pl .17 , fig. 3). In the inner exozone, thinning of the zooecial wall near interzooidal pores commonly produces moniliform profiles ( Pl .17 , figs. 1, 5c). In the outer exozone, interzooidal pores are almost cylindrical with little flare ( Pl . 17, fig. 6b; Pl. 19, fig. 1), resulting in nearly parallel-sided zooecial walls. Slight pustulose thickenings are sometimes seen in the outer exozone (Pl. 16, fig. 1g). Cross sections of zooccial chambers generally are smoothly rounded and subcircular to elliptical ( Pl .19 , fig. 1).
TABLE II
STATISTICAL SUMMARY OF MEASUREMENTS OF DIPLOCAVA INCONDITA CANU AND BASSLER

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} \mathrm{Zr}_{\mathrm{r}}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zoarial } \\ & \text { Br-CsSn-MxDn } \end{aligned}$ | 5. | 3. |  |  |  |  |  | 5 | 1 |
| $\begin{aligned} & \text { Zooecial-Exozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 28 | . 05 | . 20 | . 04 | 20 | 139 | 139 | 5 | 2 |
| Zche-CsSn-NMxDn | . 26 | . 04 | . 17 | . 04 | 22 | 139 | 139 | 5 | 2 |
| $\frac{7 c \mathrm{Ch}-\mathrm{CsSn}-\mathrm{MxDn}}{Z \mathrm{cCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 1.7 | 1.0 | 1.2 | . 1 | 11 | 139 | 139 | 5 | 2 |
| CdZcWl-Th | . 146 | . 015 | . 068 | . 026 | 38 | 139 | 139 | 5 | 2 |
| ZdPr-MnDr | 5. | 0. |  |  |  | 138 | 138 | 5 | 3 |
| Simple External Wall SEW-Th | . 061 | . 016 | . 036 | . 011 | 31 | 33 | 33 | 4 | 4 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF DIPLOCAVA INCONDITA CANU AND BASSLER

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Simple External Wall SEW-CsSn-MxDn | . 093 | . 056 | . 072 | . 022 | 29 | 14 | 14 |  | 6 |
| SEW-Psdp-DPr-Cssn-MxDn | . 021 | . 008 | . 011 | . 002 | 22 | 40 | 8 | 3 | 5 |
| Diaphragms Intd-Intrvl | . 008 |  |  |  |  | 1 | 1 | 1 | 8 |
| BslD-Th | . 013 | . 002 |  |  |  | 4 | 4 | 2 | 7 |

[^2]

Text-figure 9 A-D. Histograms and cumulative curve from the lectoiype and four topotypes of Diplocava incondita Canu and Bassler. A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzoidal pores per zooecial cross section.

Simple External Walls - Simple external walls (Pl. 16, figs. 1d, g; Pl. 17, figs. 5b, c, 6a, b; Pl. 18, figs. 1, 2, 3; Pl. 19, fig. 3) were observed in most zooecia subjacent to intrazoarial overgrowths. The walls are thick, and apertures generally located in the disto-central or central part of the diaphragm. A deposit of laminated calcareous tissue, continuous with the zooecial lining, lines the aboral surface of these diaphragms sealed by overgrowths ( Pl .16 , fig. 1g; Pl. 17, fig. 6 a; Pl. 18, figs. 1, 2) and extends orally to line the peristome.

Diaphragms - Intermediate diaphragms occur rarely and, when seen, are just aboral to the aperture ( Pl .19 , fig. 2). The diaphragms are thick, laminate, and have prominent aborally flexed abutments.

Basal diaphragms occur rarely. The diaphragms are thin and nearly planar, and flex orally at the juncture with the zooecial wall to merge with the zooecial lining (Pl. 16, fig. 1 g ; Pl. 18, fig. 2).

Discussion. - Three of the four original specimens figured by Canu and Bassler (1926, pl. 10, fig. 5) were attached to a small card bearing the label "Diplocava incondita, Cotypes, USNM 69925". Two of the specimens were thin-sectioned. USNM 69925-2 was judged to be conspecific with unfigured (?) thin-sections prepared by Bassler and labeled as types by him. In addition, USNM 69925-2 was judged to be conspecific with sectioned non-type specimens from the type locality identified by Bassler as D. incondita.

The second specimen sectioned, USNM 69925-1, is a thin-walled encrusting bryozoan referable to a different, probably unnamed,

TABLE 12
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN DIPLOCAVA INCONDITA CANU AND BASSLER

Circ. Ellip. Ov. Pyr. Polyg. Triang. Irreg.

| Endozone <br> Regular | 2 | 39 | 6 |  |
| :--- | ---: | ---: | ---: | :--- |
| Sub | 10 | 66 | 6 | 2 |
| Irregular | 4 | 3 | 1 |  |
| Exozone <br> Regular |  | 6 | 2 |  |
| Sub | 5 | 4 |  |  |
| Irregular |  |  |  |  |

genus. This was the only specimen of Canu and Bassler's original suite in which a brood chamber was observed. Thus, Canu and Bassler's remarks concerning the occurrence and morphology of brood chambers are not applicable to the concept of $D$. incondita.

Genus DITAXIA von Hagenow, 1851
Type species: Ceriopora anomalopora Goldfuss, 1826, by subsequent designation, D'Orbigny (1854, p. 952).
1826. Pars Ceriopora Goldfuss, Petrefacta Germaniae, vol. 1, p. 33.
1830. Pars Heteropora Blainville, Zoophytes, Dictionnaire de Science Naturelles, vol. 60, p. 382.
1834. Pars Heteropora Blainville, Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 417.
1851. Ditaxia von Hagenow, Die Bryozoen der Mastrichter Kreidebildung, Cassel, p. 49.
1854. Ditaxia von Hagenow, D’Orbigny, Terrain Crétacé Bryozoaires, Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 953.
1881. Polytaxia Hamm, Inaugural-Dissertation zur Erlangung der Doctorwurde Von der Philosophischen Facultät der Friedrich-Wilhelms-Universität zu Berlin, p. 41.
1899. Ditaxia von Hagenow, Gregory, Catalogue of fossil Bryozoa in Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 1, p. 406.
1922. Ditaxia von Hagenow, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 101.
1953. Ditaxia von Hagenow, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G72.

Tentative diagnosis. - Zoaria encrusting and branching. Branches bifoliate and commonly frondose, less commonly subcylindrical. Endozones and exozones strongly differentiated and intergrade in narrow zone of zooecial flexure. Zooecia dimorphic. Large dimorphs occurring in intermonticular areas; small dimorphs occurring in intermonticular and monticular areas.

In exozone, large dimorphs have lunaria-like structures composed of subgranular calcite. Lunaria-like structure becomes obscure near aperture, merging with indistinctly laminate tissue. Thin zooecial lining commonly present.

Small polymorphs have discontinuous patches of homogeneous to subgranular calcite in cortex, alternating with indistinctly laminate tissue. Laminae crenulate, broadly curved convex orally, and sometimes continuous across zooecial boundary zone.

Intermediate diaphragms occur rarely.
Taxa included. - Only the type species, D. anomalopora; in-
ternal characters of other species assigned to Ditaxia are presently unknown to the author.

Discussion. - In the large zooecia of Ditaxia, most of the cortex of the proximal wall is composed of calcareous tissue which is distinctly different from calcareous tissue in the remainder of the wall. This light-colored, homogeneous tissue is called a lunaria-like structure here, and is inferred to have been originally granular because remnants of originally laminate tissue are preserved elsewhere in the zooecial walls. Recently, studies of lunarial microstructure were made in some Paleozoic cystoporates by Utgaard (1968a, b), and in post-Paleozoic lichenoporids by Brood (1970b). Unfortunately, comparison of lunaria in these taxa to the lunaria-like structure in Ditaxia cannot be made because microstructure is poorly preserved in the specimens of Ditaxia which are available for study.

The recognition of a lunaria-like structure poses some questions concerning continued assignment of Ditaxia to the cerioporids. On the basis of mode of growth, wall structure, and diaphragms, Ditaxia is not readily assignable to other taxa as presently understood, e.g., lichenoporids and is provisionally retained in the cerioporids.
1826. Ceriopora anomalopora Goldfuss, Petrefacta Germaniae, vol. 1, p. 33, pl. 10, figs. 5c, d.
1830. Heteropora anomalopora (Goldfuss), Blainville, Zoophytes, Dictionnaire de Science Naturelles, vol. 60, p. 382.
1834. Hetcropora anomalopora (Goldfuss), Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 417.
1851. Ditaxia anomalopora (Goldfuss), Von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, Cassel, p. 49, pl. 4, fig. 9c.
1881. Polytaxia anomalopora (Goldfuss), Hamm, Inaugural-Dissertation zur Erlangung der Doctorwurde Von der Philosophischen Facultät der Friedrich-Wilhelms-Universität zu Berlin, p. 41.
1899. Ditaxia anomalopora (Goldfuss), Gregory, Catalogue of Fossil Bryozoa in Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 1, p. 406.
1922. Ditaxia anomalopora (Goldfuss), Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 101.

Type. - Specimen UB 119 is designated as the lectotype. The lectotype was figured by Goldfuss (1826, pl. 10, figs. 5c, d) and by Von Hagenow (1851, pl. 4, fig. 9c).

Type locality and horizon. - Goldfuss (1826, p. 33) cited the
locality as "Petersberge bei Maastricht". Rocks exposed at this locality are Maastrichtian in age. Therefore, the horizon is assumed to be Cretaceous, Maestrichtian.

Material studied. - Four thin-sections and four acetate peel replicas were made from the lectotype UB 119; most of the lectotype remained after sectioning. In addition, eight topotypes in the collections of the National Museum of Natural History were thin-sectioned: four specimens labeled "USNM Loc. 2405, Up. Cret.-Maastr., Geulem, Maastricht, Netherlands"; and three specimens labeled "USNM Loc. 2965A, Cret.-Maastr., Maastricht, Netherlands". Duplicate acetate peels of specimens thin-sectioned are in the Bonn collection and the National Museum of Natural History collection. The author has duplicate peels of the lectotype and the specimens from USNM Loc. 2405.

Description. -
Median layer - The median layer is thick (Table 13, MdnLyrTh) and laminate. In cross section, the laminae form steep Vshaped configurations pointing proximally. The boundary zone of the median layer is commonly marked by a distinct, dark line (Pl. 20, fig. 1e).

Endozone - In the recumbent endozone, the zooecial chambers have trianglar cross sections, and become polygonal distally (Pl. 20, fig. 1e). Zooecial growth axes are nearly straight throughout the endozone, diverging from the distal growth axis at a low angle, commonly less than $30^{\circ}$ ( Pl. 20, fig. 1d). Zooecial walls are thin and parallel-sided with a homogenous to subgranular cortex, and a thin, dark-colored zooecial lining except on the recumbent wall (median layer).

Exozone - In the exozone, the large dimorphs are irregular in distribution (Pl. 20, figs. 1a, b). The walls of adjacent large and small dimorphs are subsymmetrically to nonsymmetrically thickened across the boundary zone. The walls of large dimorphs are commonly nearly parallel-sided. Zooecial chambers have subcircular, elliptical, or subelliptical cross sections. The walls of large dimorphs protrude slightly above the zoarial surface.

Small dimorphs are either scattered between large dimorphs in monticular areas, or are concentrated in these areas. Monticular
TABLE 13
STATISTICAL SUMMARY OF MEASUREMENTS OF DITAXIA ANOMALOPORA (GOLDFUSS)

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF DITAXIA ANOMALOPORA (GOLDFUSS)

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF DITAXIA ANOMALOPORA (GOLDFUSS)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zooecial-Small Polymorphs - Exozone ZcCh-CsSn-MxDn | . 10 | . 03 | . 07 | . 02 | 25 | 57 | 57 | 1 | 1 |
| Zch-CsSn-NMxDn | . 08 | . 02 | . 06 | . 01 | 23 | 57 | 57 | 1 | 1 |
| $\frac{\text { ZcCh-CsSn-MxDn }}{\text { ZcCh-CsSn-NMxDn }}$ | 1.9 | 1.0 | 1.3 | . 2 | 15 | 57 | 57 | 1 | 1 |
| CdZcWl-Th | . 147 | . 023 | . 064 | . 028 | 43 | 57 | 57 | 1 | 1 |
| Zooecial - Small Polymorphs - Endozone $\mathrm{CnZcCh}-\mathrm{CsSn}-\mathrm{MxDn}$ | $1 .$ | 0. |  |  |  | 57 | 57 | 1 | 1 |

[^3]areas are patchlike in distribution ( Pl .20 , fig. 1b). The walls of small dimorphs commonly show annular thickenings. The thickenings of adjacent small dimorphs are symmetrical across the zooecial boundary zone. Monilar profiles are circular, obovate to inverse pyriform, or less commonly, nonsymmetrically thickened across the boundary zone. The chambers of small dimorphs are commonly elliptical in cross section.

Diaphragms - Intermediate diaphragms were sometimes observed. The diaphragms are thin ( .005 to .008 mm ) and planar, and were most commonly observed close to the zooecial aperture, although they were sometimes seen as deep as the zone of zooecial flexure.

## TABLE 14

FREQUENCY OF VISUALLY ESTIMATED OUTLINE OF ZOOECIAL CHAMBERS IN DITAXIA ANOMALOPORA (GOLDFUSS)*

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Exozone - Large Polymorphs |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |
| Sub | 10 | 7 | 1 |  |  |  |  |
| Irregular |  | 1 |  |  |  |  |  |
| Exozone - Small Polymorphs |  |  |  |  |  |  |  |
| Regular | 4 | 29 | 5 |  |  |  |  |
| Sub | 8 | 6 | 1 |  |  |  |  |
| Irregular |  | 4 |  |  |  |  |  |
| *Outlines of zooecial chambers were estimated on the lectotype, UB 120 (50 zooecia) and USNM 2405-2 (50 zooecia). |  |  |  |  |  |  |  |

Discussion. - The designation of lectotype is consistent with a previous restriction made by Von Hagenow (1851). Goldfuss figured three specimens in his plate 10, figure 5, as Ceriopora anomalopora. Von Hagenow considered only the specimen figured in plate 10 , figures 5 c , d, as $D$. anomalopora, and refigured this specimen in his plate 4, figure 9c. This restriction was later followed by Gregory (1899, p. 406) and Canu and Bassler (1922, p. 101). The word "Holotypus" is written on a label with the specimen, but the label was most certainly added much later than 1826 by someone other than Goldfuss.

Remarks on dimorphism. - Zooecial dimorphism in the exozone is expressed in the bimodal frequency distribution of the zooecial


Text-figure 10 A-D. Histograms and cumulative curve from the lectotype and one topotype of Ditaxia anomalopora (Goldfuss). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.
diameters (Table 13A), and in the appearance and apparent structure (see below) of the zooecial walls. The general terms, large and small polymorphs, are used for typical members of each dimorph rather than autozooecia, and the like because the characters described are not here considered sufficient to infer function or description of enclosed soft parts.

Remarks on microstructure. - Most of the specimens studied, including the lectotype, show poorly preserved microstructure, and in only three specimens were structures preserved well enough to attempt interpretation. The patchy occurrence of nondescript, calcareous material, and the irregular occurrence of a laminar tissue may well be, in part, artifactual and due to diagenetic changes of primary structures. The appearance of the wall in some large polymorphs is strongly suggestive of lunaria-like deposits. These structures, however, are not here identified as lunaria because of four factors:

1) The structures identified as possible lunarial deposits, although commonly occurring on the proximal side of the zooecium, were sometimes observed in other positions around the zooecial cavities of large polymorphs. Lunarial deposits, typically, are emplaced in the proximal portion of a zooecial wall (Utgaard, 1968a, p. 1033).
2) As seen in tangential section, patches of calcareous tissue, similar in appearance to the possible lunarial structures, were seen in zooecial walls of small polymorphs.
3) The possible lunarial deposits were not associated with any lunaria-like inflections of the outline of the zooecial cavity.
4) In well-preserved fistuliporoids, ceramoporoids, or lichenoporids, the lunaria project orally from the zoarial surface as hoods. These were not seen on any specimens of $D$. anomalopora.
The lunaria-like structure is often missing, or poorly expressed, in shallow tangential section. Sometimes, as seen in longitudinal section, the lunaria-like deposit is capped by indistinct laminate tissue. Possibly the mode of growth changes slightly in the outer exozone, and lunaria-like tissue is not deposited, suggesting ontogenetic control.

## Genus HAPLOECIA Gregory, 1896

Type species: Millepora straminea Phillips, 1829, by original designation.
1896. Haploccia Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, p. 157.
1922. Haploccia Gregory, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, pp. 97-98.
1953. Haploecia Gregory, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G71.
1965. Dendroccia Cotillon, and Walter, Soc. Géol. France, Bull., vol. 7, ser. 7, pp. 934-935.

Tentative diagnosis. - Zoaria branched; distally growing branches intersect and anastomose. Branches have coaxial endozones and exozones. Zooecia intersect zoarial surface obliquely. Branches commonly encrusted by one or more intrazoarial overgrowths.

In endozone, zooecia with smaller diameters often surround single large zooecium growing parallel to distal growth axis of branch and located close to center of branch.

In exozone, cortex composed of light-colored, indistinctly laminated, calcareous tissue. Laminae slightly crenulate and, in general, broadly arched, convex orally. Lamination commonly becomes obscure or disappears in outermost cortex. Zooecial linings dark-colored, with crenulate, longitudinally directed laminae. Lateral spinelike extension of light-colored cortex tissue sometimes extends into zooecial chamber as mural spines, or sometimes submerged beneath thick deposits of zooecial lining.

All zooecia distal to unencrusted growing tips sealed by simple external walls. Intermediate and basal diaphragms occur but not common.

Taxa included. - Based on the examination of primary types, two species are here considered to be correctly assigned to Haploecia: Ceriocava multilamellosa Canu and Bassler, 1922 [Lower Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland], and the type species, Millepora straminea Phillips, 1829 (Middle Jurassic, Yorkshire, England). Internal characters of other species assigned to Haploecia are unknown to me.

Discussion. - On the basis of wall structure and occurrence of diaphragms, Haploecia is retained in the cerioporids. The occurrence
of simple external walls and attendant modification of growth habit is known to occur in only one other cerioporid genus, Diplocava. Simple external walls resemble externally the porous, foraminate, apertural seals seen in the Salpingina. Viskova has recently studied genera assignable to the Salpingina and has redefined three families on the basis of internal characters (Viskova, 1965, 1968). Assignment of Haploecia would be consistent with the revised concept of the Eleidae d'Orbigny except for differences in wall structure. Viskova (1968, p. 175) reported that zooecial walls in eleids are longitudinally fibrous. In specimens of Haploecia available for study, wall structure was poorly preserved. Orally convex lineations (Pl. 23, fig. 1c; Pl. 26, figs. 1b, c; Pl. 28, fig. 3b ), however, are interpreted as remnants of original lamination.

Gregory (1896, p. 157) allied Haploecia with the entalophorids, and in 1899 (pp. 288-9) suggested that Haploecia was an early stage in the evolution of the Eleidae from the Entalophoridae. The melicerititids, a later evolutionary stage in Gregory's scheme, are characterized by tubular zooecia, but are reputed to possess both opercular and avicularian structures (Gregory, 1899, pp. 287-292; Levinsen, 1912, pp. 10-13, 19; Bassler, 1953, p. G75). The occurrence of structures presumably diagnostic for cheilostomes (Bassler, 1953, p. G147) in tubular bryozoans led Gregory (1899, p. 287) to state that "The Cretaceous family, the Eleidae, is important, as it breaks down the distinction between the Cheilostomata and Cyclostomata on these characters".

The external similarity of the type species of Haploecia to some Cheilostomata is striking (Pl. 23, fig. 1b). As observed by Gregory (1899, p. 288), " . . the zooecia, at their distal ends, are hexagonal, bounded by ridges, and have a small subterminal aperture at the upper part. This arrangement is very similar to that of the genus Cellaria." This similarity, however, is due to convergence, as can be ascertained by reference to internal structures. The apertural structure resembling a porous frontal wall is a simple external wall. The zooecia in Haploecia are tubular, rather than boxlike; and the zooecial walls in Haploecia are compound, being shared with adjacent zooecia rather than the cuticle-lined individual cases typical of the Cheilostomata.

Canu and Bassler (1922, pp. 97-8) were apparently the first to study thin-sections of specimens assignable to the type species. They observed the enlarged axial zooecium, the simple external walls ("facettes"), the absence of mesopores, and the general absence of other diaphragms. They referred Haploecia to the Ceriocavidae.

As presently understood, Ceriocava and Haploecia exhibit large morphologic differences, although the mode of growth is similar. Both have coaxially arranged endozones and exozones, and show increased diameters of zooecial chambers from endozone to exozone. In Ceriocava, however, zooecia apparently continued to grow in the exozone for much of the life of the colony; in Haploecia, orally directed growth terminates relatively soon after zooecia become exozonal in character. Terminal diaphragms are emplaced in Ceriocava, simple external walls in Haploecia. Basal diaphragms are numerous and closely spaced in Ceriocava but were rarely emplaced in Haploecia. Anastomosis occurs in Ceriocava but is associated with the bifurcation of two branches rather than with the intersection of their growing tips as seen in Haploecia.

Cotillon and Walter (1965, pp. 934-5) erected Dendroecia because of the nature of the budding and the appearance of the ovicells in specimens they assigned to Ceriocava multilamellosa (Canu and Bassler, 1922). The internal morphology of the primary types of Ceriocava multilamellosa is consistent with the generic concepts of Haploecia, and Dendroecia is determined to be a junior subjective synonym.

Haploecia straminea (Phillips), 1829 Pl. 23, figs. 1a-f; Pl. 24, figs. 1a-f; Pl. 25, figs. 1a-d, 2a-b; Pl. 26, figs. 1a-d
1829. Millcpora straminea Phillips, Illustrations of the Geology of Yorkshire. Description of the strata and organic remains of the Yorkshire Coast, pp. 144, 149, pl. ix, fig. 1.
1893. Pustulopora straminea (Phillips), Gregory, Yorkshire Philos. Soc., Ann. Rept., p. 60, text-fig. 2.
1896. Haploccia straminea (Phillips), Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, pp. 159-161, text-fig. 12, p. 160.
1922. Haploecia straminea (Phillips), Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, pp. 97-8, pl. 14, figs. 14, 15, text-fig. 25, p. 98.
1945. Haploecia straminea (Phillips), Melmore, Catalogue of Types and Figured Specimens in Geological Department of Yorkshire Museum, p. 216.

Type. - The lectotype is YM-T81/2 (figured by Gregory,

1893, text-fig. 2, p. 60, and 1896, text-fig. 12, p. 160) by indication, Gregory (1896, explanation of text-fig. 12, p. 160).

Type locality and horizon. - The label with the type specimens bears only information bearing on Gregory's publication, and no locality data are given. Phillips (1829, pp. 144, 149) cited the localities as "Scarborough (very rare), from the Cornbrash; Gristhorpe, Cloughton, Owlton, Crambe, Westow, Ellerker, from Gray Limestone or Bath oolite, England." Sediments bearing fragments of bryozoans identified with $H$. straminea exposed at these localities are questionably referred to the Bajocien (pers. comm. J. W. Neale).

Material studied. - The lectotype YM-T81/2 and paralectotype YM-T81/1 were borrowed from the Yorkshire Museum. Thin-sections and acetate peels were made from both the lectotype and paralectotype.

In addition, an encrusted specimen of H. straminea was revealed in a thin section of Ceriocava corymbosa (Lamouroux) [USNM 32181-2, from the Bathonien, Langrum (Calvados), France]. Acetate peel replicas of all specimens sectioned are preserved in the collections of the National Musum of Natural History and of the author.

Description. -
Growth habit - Zoaria are profusely branched with branches growing distally in all directions (Pl. 23, fig. 1a). Branches intersect and anastomose in the proximal portion of the colony forming a densely matted network. Branches have subcircular to elliptical cross sections and wedge-shaped to rounded growing tips.

Endozone - Zooecial walls are thin and slightly undulatory (PI. 24, fig. 1c). Generally, the walls are nearly parallel-sided, but prominent asymmetrical thickenings are sometimes observed. The walls are light-colored and granular. Zooecial linings are thin or missing in the inner endozone ( Pl .23 , figs. 1c, d, f; Pl. 24, fig. 1c). Interzooidal pores are rarely seen.

Exozone - Near the tips of unencrusted primary branches, zooecial walls are sometimes thickened asymmetrically across the zooecial boundary zone (Pl. 23, fig. 1d). Walls have submoniliform profiles because of moderate variation in thickness longitudinally (Pl. 23, fig. 1d) and because of the relatively wide flare of interzooidal pores (Pl. 23, figs. 1c, d, e). The outer exozone portions of
zooecial walls are thicker in a gradient directed proximally from unencrusted primary branch tips (Table 15) in at least some encrusted primary branch tips (Pl. 24, fig. 1a), and in intrazoarial overgrowths (Pl. 24, fig. 1b). Thicker zooecial walls generally are nearly parallel-sided because of in-filling of thinner portions of the wall, and because of reduction in the "flare" of interzooidal pores (Pl. 26, figs. 1c, d). Zooecial cross sections are generally subelliptical (Table 16) except at apertural level. At the apertural level, the wall thins rapidly to a rimlike extension (Pl. 23, fig. 1c; Pl. 25, fig. 2b; Pl. 26, fig. 1b), and zooecia have polygonal, commonly hexagonal outlines (Pl. 23, fig. 1b; Gregory, 1896, text-fig. 2, p. 60).

Simple External Walls - External walls were observed in nearly all zooecia more than about 1 mm from unencrusted branch tips. The exterior walls are thicker and generally appear more robust in a gradient directed proximally from unencrusted branch tips (compare Pl. 23, figs. 1c, d, and Pl. 25, figs. 2a, b, to Pl. 24, figs. 1c, f and Pl. 26, figs. 1a-d). Peristomes are low near unencrusted branch tips (Pl. 23, fig. 1b); but proximally from branch tips, peristomes may extend as much as .2 mm above the zoarial surface (Pl. 24, fig. If; Pl. 26, figs. 1b, c, d). Peristomes have circular to elliptical cross sections and are located in the centro-distal portion of each zooecium (Pl. 24, figs. 1e, f). Measurements of the maximum dimensions of peristomial cross section (the restricted skeletal aperture) and cross sections of diaphragmal pores are summarized in Table 15.

Diaphragms - Both intermediate and basal diaphragms (Pl. 25, fig. 1b) occur but, in general, are not numerous. Diaphragms emplaced in the almost wholly recumbent zooccia of some interzoarial overgrowths are attached, in part, to simple external walls (Pl. 26, fig. 1b). In addition, basal diaphragms were sometimes observed in peristomes (Pl. 26, fig. 1b).

Discussion. - The original figure by Phillips (1829, p. 19) is generalized and of little use in identifying other specimens with $H$. straminea. As noted by Gregory (1896, p. 160). "This species was figured so imperfectly by Phillips that it had been variously interpreted by foreign authors". Gregory refigured specimen T81/2 twice (1893, text-fig. 2, p. 60; 1896, text-fig. 12, p. 160); both figures are somewhat idealized illustrations showing regularly arranged hexa-
TABLE 15
STATISTICAL SUMMARY OF MEASUREMENTS OF HAPLOECIA STRAMINEA (PHILLIPS)

STATISTICAL SUMMARY OF MEASUREMENTS OF HAPLOECIA STRAMINEA (PHILLIPS)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | N Zr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zooecial-Exozone }{ }_{1} \text { (con.) } \\ & \text { ZcWlLn-Th }^{\text {O }} \end{aligned}$ | . 005 | . 000 | . 003 | . 002 | 46 | 17 | 17 | 1 | 1 |
| ZdPr-MnDr | . 008 | . 003 | . 005 | . 002 | 37 | 16 | 16 | 1 | 1 |
| $\begin{gathered} \text { Zooecial }^{\text {ECChozone }_{2}} \\ \text { CCh-CsSn-MxDn } \end{gathered}$ | . 20 | . 16 | . 18 | . 02 | 13 | 8 | 8 | 1 | 1 |
| ZcCh-CsSn-NMxDn | . 14 | . 08 | . 10 | . 02 | 18 | 8 | 8 | 1 | 1 |
| $\frac{Z \mathrm{cCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 2.1 | 1.5 | 1.7 | . 2 | 11 | 8 | 8 | 1 | 1 |
| $\mathrm{CdZcWl}-\mathrm{Th}$ | . 13 | . 06 | . 10 | . 02 | 24 | 8 | 8 | 1 | 1 |

STATISTICAL SUMMARY OF MEASUREMENTS OF HAPLOECIA STRAMINEA (PHILLIPS)

| Character | O.R.* |  | X* | $S^{*}$ | C. V. | N | NZc | $\mathrm{N} Z \mathrm{r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zooecial-Exozone } 2 \text { (con.) } \\ & \text { ZdPr-Cn } / \mathrm{ZcCsSn} \end{aligned}$ | 2. | 0. |  |  |  | 8 | 8 | 1 | 1 |
| ZcWlLn-Th | . 035 | . 012 | . 024 | . 007 | 32 | 8 | 8 | 1 | 1 |
| $\begin{aligned} & \text { Zooecial-Endozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 17 | . 02 | . 10 | . 04 | 39 | 30 | 30 | 1 | 1 |
| Zch-CsSn-NMxDn | . 12 | . 01 | . 07 | . 02 | 39 | 30 | 30 | 1 | 1 |
| $\frac{\text { ZcCh-CsSn-MxDn }}{\text { ZcCh-CsSn-NMxDn }}$ | 2.4 | 1.0 | 1.6 | . 3 | 21 | 30 | 30 | 1 | 1 |
| CdZcWl-Th | . 03 | . 01 | . 01 | . 005 | 39 | 30 | 30 | 1 | 1 |
| ZdPr-Cn/ZcCssn | 1. | 0. |  |  |  | 30 | 30 | 1 | 1 |
| Simple External Wall SEW-CsSn-MxDn | . 077 | . 058 | . 067 | . 022 | 36 | 13 | 13 | 2 | 2 |
| SEW-Psdp-CsSn-MxDn | . 008 | . 003 | . 005 | . 002 | 37 | 16 | 16 | 1 | 1 |
| Diaphragms Bsld-Th | . 006 | . 001 |  |  |  | 7 | 7 |  | 2 |
| *In millimeters <br> ${ }_{2}$ Encrusted branch tips, or more <br> 1. Lectotype: YM T81/2. <br> 2. Lectotype and paralectotype <br> 3. Paralectotype: YM T81/1. | $m$ from <br> /1. | ncruste <br> Y TO | anch <br> CIME |  |  |  |  |  |  |



Text-figure 11 A-D. Histograms and cumulative curve from the lectotype and paralectotype of Haploecia straminea (Phillips). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores zooecial cross section.


Text-figure $12 \mathrm{~A}-\mathrm{C}$. Histograms showing variation in thickness of peristomial diaphragms in Haploccia straminea (Phillips). A. More than 3.0 mm from tip of encrusted branch in YM T81/1. B. More than 1.2 mm , less than 3.0 mm from branch tip in YM T81/2. C. Less than 1.2 mm from tip of unencrusted branch in YM T81/2.

TABLE 16
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN HAPLOECIA STRAMINEA (PHILLIPS)

Circ. Ellip. Ov. Pyr. Polyg. Triang. Irreg.

| Exozone <br> Regular | 1 | 1 | 1 |  |  |
| :--- | ---: | ---: | ---: | ---: | :--- |
| Sub | 11 |  | 2 |  |  |
| Irregular | 1 |  |  |  |  |
| Endozone <br> Regular | 1 | 4 |  |  |  |
| Sub | 11 | 4 | 1 | 4 |  |
| Irregular | 3 | 1 |  |  | 1 |

gonal zooecial apertures sealed by simple external walls. In the explanation, Gregory said that the figure is, "Part of the type specimen of Haploccia straminea (Phil.)".

Haploecia multilamellosa (Canu and Bassler), 1926 Pl. 27, figs. 1a-e, 2a, b, 3a-d; Pl. 28, figs. 1a-e, 2, 3a, b; Pl. 29, figs. 1a-e, 2a-c, 3; Pl. 30, figs. 1a, b, 2, 3; Pl. 31, figs. 1a, b, 2a, b
1926. Ceriocava multilamellosa Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 67, pp. 68-70, text-fig. 35A-F, p. 69, pl. 9, figs. 1-10.
1965. Dendroecia multilamellosa (Canu and Bassler), Cotillon, and Walter, Soc. Géol. France, Bull., ser. 7, vol. 7, pp. 934-5.

Type.-USNM 69922-1 is designated as the lectotype. This specimen was figured by Canu and Bassler (1926, pl. 9, figs. 1 middle specimen, 5,6 ), and here (Pl. 27, figs. 1a-e).

Type locality and horizon. - Lower Cretaceous (Valangian): Sainte Croix (Vaud), Switzerland.

Material studied. - Only the lectotype and single paralectotype of the five specimens originally figured by Canu and Bassler (1926, pl. 9, figs. 1-10) have been found in the National Museum of Natural History collection. Both specimens bear the label "cotype." The lectotype, USNM 69922-1, is the specimen figured by Canu and Bassler (pl. 9, figs. 1 - the middle specimen, 5, 6). Three thinsections and four peels on one acetate slide were prepared from the lectotype. Paralectotype specimen USNM 69922-2 is the specimen figured in Canu and Bassler (1926, pl. 9, figs. 1 - second specimen from the right, 7 and 8). Four unnumbered thin-sections, labeled type and prepared by R. S. Bassler, are considered paralectotypes.

The thin-sections were figured by Canu and Bassler (text-fig. 35, p. 69) as follows: No. 41, text-fig. 35B, C; No. 41.5, text-fig. 35D; No. 41.7, text-fig. 35E; No. 41.6, text-fig. 35F. Thin-sections and acetate peels were made from nine topotypes identified by R. S. Bassler. These bear the number USNM Loc. 2384.

## Description. -

Growth habit - Branches are roughly cylindrical, consisting of a primary branch with a well-defined coaxial endozone and up to three thin (Table 17, Ov-Th) intrazoarial overgrowths. Zooecial apertures are generally arrayed in parallel ranges. Zooecia in adjacent ranges commonly alternate in position longitudinally (Pl. 27, figs. 1b, 3b, d; Pl. 28, fig. 3a; Pl. 29, fig. 1b).

Distally growing branches sometimes intersect and anastomose (Pl. 29, figs. 1a, 2a). Where the plane of intersection is nearly normal to the direction of zooecial growth, each zooecium is generally sealed by a thin, nonporous diaphragm (Pl. 29, figs. 1c, d, 2b, lower center, $2 c)$. Where the plane of intersection is oblique to zooecial growth, the zooecial walls of intersecting zooecia merge and continue to grow orally ( Pl . 29, figs. 2a, b, upper center).

Endozone - Zooecial walls are parallel-sided and occasionally have rounded to spinose projections (Pl. 28, fig. 1d). Longitudinally, the walls are moderately undulatory (Pl. 28, figs. 1a, d; Pl. 29, fig. 1c).

Exozone - Zooecial walls have submoniliform profiles and are symmetrical across the zooecial boundary zone. Zooecial cross sections are commonly elliptical. In the outer exozone, mural spines are sometimes numerous within individual zooecia. The spines have light-colored (subgranular) cores which contrast with the darkcolored laminated tissue of the zooecial lining (Pl. 28, figs. 1c, d; Pl. 31, fig. 2b). Interzooidal pores are nearly cylindrical and only slightly flared.

Simple External Walls - External walls were observed in all zooecia subjacent to intrazoarial overgrowths. The walls are relatively thin and composed of dark-colored, wavy, laminate tissue similar in appearance to the zooecial lining (Pl. 30, figs. 1a, b). Laminae were sometimes observed to flex aborally and merge with the zooecial lining (Pl. 30, figs. 1a, b; Pl. 31, figs. 1a, b, 2b). Peristomes are commonly subcircular, located centrally, and extend only
TABLE 17
STATISTICAL SUMMARY OF MEASUREMENTS OF HAPLOECIA MULTILAMELLOSA (CANU AND BASSLER)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} Z \mathrm{r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zoarial |  |  |  |  |  |  |  |  |  |
| Br-CsSn-MxDn | 2.4 | 1.2 | 1.7 | . 4 | 24 | 7 |  | 7 | 1 |
| $\mathrm{PrBr}-\mathrm{CsSn}-\mathrm{MxDn}$ | 1.5 | . 8 | 1.0 | . 2 | 22 | 7 |  | 7 | 1 |
| $\mathrm{Ov}-\mathrm{Th}$ | . 26 | . 13 | .17 | . 09 | 55 | 14 |  | 7 | 1 |
| $\begin{aligned} & \text { Zooecial-Exozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 20 | . 10 | . 15 | . 02 | 15 | 41 | 41 | 3 | 2 |
| Zch-CsSn-NMxDn | . 11 | . 05 | . 08 | . 02 | 18 | 41 | 41 | 3 | 2 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 3.3 | 1.4 | 2.0 | . 4 | 19 | 41 | 41 | 3 | 2 |
| CdZcWl-Th-Lng | . 26 | . 08 | . 18 | . 07 | 37 | 16 | 16 |  | 3 |
| CdZcWl-Th-Trv | . 16 | . 05 | . 09 | . 03 | 36 | 15 | 15 |  | 4 |
| ZdPr-Cn/ZcCsSn | 6. | 0. |  |  |  | 41 | 41 | 3 | 2 |
| ZcWlLn-Th | . 026 | . 003 | . 013 | . 005 | 41 | 34 | 34 | 2 | 9 |
| ZcSp-Cn/Zc | 8. | 0. |  |  |  | 41 | 41 | 3 | 2 |

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF HAPLOECIA MULTILAMELLOSA (CANU AND BASSLER)

\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline Character \& \multicolumn{2}{|l|}{O.R.*} \& X* \& S* \& C. V. \& N \& NZc \& \(\mathrm{N} \mathrm{Zr}^{\text {r }}\) \& Spec. Code \\
\hline ZdPr-MnDr \& . 010 \& . 003 \& \& \& \& 3 \& 3 \& 1 \& 5 \\
\hline Zooecial-Endozone \(\mathrm{CnZch}-\mathrm{CsSn}-\mathrm{MxDn}\) \& . 16 \& . 08 \& . 11 \& . 03 \& 27 \& 7 \& 7 \& 7 \& 1 \\
\hline ZdPr-MnDr \& . 016 \& . 008 \& \& \& \& 3 \& 3 \& 1 \& 5 \\
\hline Simple External Walls SEW-Th \& 0.27 \& . 006 \& . 014 \& . 006 \& 44 \& 42 \& 42 \& 4 \& 6 \\
\hline SEW-Pst-CsSn-MxDn \& . 082 \& . 055 \& . 071 \& . 012 \& 17 \& 8 \& 8 \& 2 \& 7 \\
\hline SEW-Psdp-CsSn-MxDn \& . 013 \& . 005 \& . 008 \& . 002 \& 25 \& 20 \& 4 \& 1 \& 8 \\
\hline \multicolumn{9}{|l|}{*In millimeters} \& \\
\hline \begin{tabular}{l}
1. Lectotype USNM 69922 2384-9 (1 each). \\
2. Lectotype USNM 69922 \\
3. Lectotype USNM 69922 \\
4. Lectotype USNM 69922 \\
5. Topotype identified by \\
6. Lectotype USNM 69922 \\
7. Topotypes identified b \\
8. Topotype identified by \\
9. Lectotype USNM 69922
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4-7,2384-8,
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\end{tabular}

slightly above the zoarial surface. Peristomial apertures subjacent to overgrowths are commonly sealed by dark-colored laminate tissue continuous with the diaphragm (Pl. 31, figs. 1b, 2a).

Diaphragms - Intermediate diaphragms (Pl. 30, fig. 3) were observed rarely in primary branches. The diaphragms are planar to strongly convex orally and merge continuously with the zooecial lining. Diaphragms are moderately thin (Table 17, IntD-Th).

Intrazoarial overgrowths - Zooecia bud obliquely to the basal layer and intersect the zoarial surface obliquely; commonly, little or no zooecial bend is seen (Pl. 28, fig. 1a; Pl. 29, figs. 1a, c). Although zooecia are relatively short, both thin-walled (endozonal) and thick-walled (exozonal) portions are present. Mural spines sometimes project from the basal layer (Pl. 30, fig. 1b; Pl. 31, fig. 2b). Intermediate diaphragms were rare to moderately numerous. At the most, each zooecium contained a single diaphragm.

Discussion. - Canu and Bassler, in their original description of Ceriocava multilamellosa, noted that "The appearance of facettes in the genus Ceriocava still remains a mystery." Cotillon and Walter (1965, p. 935) recognized other morphologic characteristics that were not compatible with their concept of Ceriocava based on the type species, C. corymbosa (Lamouroux). Among these differential characteristics in $C$. multilamellosa were the lack of transverse diaphragms and mesopores, zooecia less bent, increase in zoarial diameter by overgrowth, and very different appearance of the ovicell.

TABLE 18
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL
CHAMBERS IN HAPLOECIA MULTILAMELLOSA
(CANU AND BASSLER)

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Exozone <br> Regular |  | 31 | 7 |  |  |  |  |
| Sub |  | 2 |  |  |  |  |  |
| lrregular | 1 |  |  |  |  |  |  |
| Endozone <br> Regular | 1 | 12 | 4 |  |  |  |  |
| Sub | 7 | 14 | 4 |  |  |  |  |
| Irregular |  | 1 |  |  |  |  |  |




Text-figure 13 A-E. Histograms and cumulative curve from the lectotype and two topotypes of Haploecia multilamellosa (Canu and Bassler). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness as measured in longitudinal section. D. Compound zooecial wall thickness as measured in transverse section. E. Count of interzooidal pores per zooecial cross section.

Cotillon and Walter described the appearance of ovicells as seen in specimens collected from the calcaire lumachelliques (Hauterivien and Barremien) from localities in the Basses-Alpes, France, which they assigned to C. multilamellosa. On the basis of these differences in morphology, Cotillon and Walter erected the new genus Dendroecia for C. multilamellosa Canu and Bassler.

Based on an examination of thin-sections of syntypes and topotypes identified by Canu and Bassler, C. multilamellosa has characters consistent with the tentative diagnosis of Haploecia and is assigned to Haploecia.
H. multilamellosa differs from the type species, $H$. straminea, in having branches with smaller diameters and in the more regular arrangement of zooecial apertures. Zooecia have smaller cross sections and fewer interzooidal pores per zooecial cross section, but more mural spines. Peristomes commonly are located centrally with relation to the zooecial walls. In $H$. multilamellosa, zooecia in overgrowths generally grew obliquely away from the basal layer; in the type species, zooecia usually grew recumbent to the basal layer for much of their length.

## Genus HETEROPORA Blainville, 1830

Type species: Ceriopora cryptopora Goldfuss, 1826, by subsequent designation, Gregory (1896, p. 201).
1830. Heteropora Blainville, Zoophytes: Dictionnaire de Science Naturelles, vol. 60, p. 381.
1834. Heteropora Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 417.
1851. Pars Ceriopora Goldfuss, Von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, p. 53.
1896. Heteropora Blainville, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, p. 201.
1909. Heteropora Blainville, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 185.
1933. Non Heteropora Blainville, Borg, Zool. Bidrag från Uppsala, Band 14, p. 255.
1953. Heteropora Blainville, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G66.

Tentative diagnosis. - Zoaria massive to branching, composed of superposed intrazoarial overgrowths. Endozones thin, directly adjacent to basal layer, or not developed.

Zooecial walls granular, becoming indistinctly laminate in outer
exozone. Laminae broadly arched, convex orally and continuous across zooecial boundary zones. Thin zooecial lining commonly present in endozone and exozone. Intermediate diaphragms common, basal diaphragms occurring rarely.

Taxa included. - Only the type species, Ceriopora cryptopora, is here included. Heteropora dichotoma, included in Heteropora by Blainville, is here considered to be correctly assigned to Ditaxia. Characters seen in thin sections of specimens assigned to $H$. dichotoma by Bassler, the third species included in Heteropora by De Blainville, are not consistent with the concept of Heteropora as understood here.

Thin sections made from topotypes of $H$. magna O'Donoghue and O'Donoghue and $H$. pacifica Borg, and sections made from a syntype and identified specimens of $H$. pelliculata Waters, revealed the following characters:

1) Zoaria have coaxial exozones and endozones.
2) Zooecia are dimorphic.
3) Zooecial walls are composed of aborally oblique laminae.
4) Terminal (and possibly peristomial), intermediate, and basal diaphragms occur.
These characters are not consistent with the concept of Heteropora, and the above species are assignable to another, probably unnamed genus.

The internal characters of numerous other species assigned to Heteropora by authors are presently unknown.

Discussion. - Heteropora was erected by Blainville for three species first described by Goldfuss (1826) and placed in his genus Ceriopora. Goldfuss (1826, p. 32) had erected the genus Ceriopora and assigned 28 species to it, all of them new. Heteropora was erected by Blainville to separate from Ceriopora Goldfuss those species which "se distingue essentiellement par l'extence de deux sortes de cellules ou de pores, les unes deux ou trois fois plus grandes que les autres . . . et composées de couches enveloppants". Blainville, in erecting Heteropora, apparently had based his diagnosis only on the figures and descriptions of Goldfuss, and had not observed a specimen assignable to the genus because he wrote, "Nous ne voudrions cependent pas assurer ce dernier point, n'ayant pas
encore analysé nous-même une espèce d'heteropore" ("ce dernier point" refers to "de couches enveloppants" Blainville, 1834, p. 417).

Gregory was the first reviser of Heteropora Blainville because in 1896 (p. 201) he cited $H$. cryptopora (Goldfuss) as the type species of Heteropora. In 1909 (p. 185), he noted that "Of these three species $H$. cryptopora was mentioned first and of the others the C. anomalopora, Goldf., is a Ditaxia and C. dichotoma, Goldf., is a Sparsicavea."

A single character, zooecial dimorphism, had been consistently cited by authors, except Haime (1854), as diagnostic for Heteropora. The occurrence of dimorphic zooecia was not, however, confirmed in this study. "Polymorphism is discontinuous variation in the morphology of zooids arising at the same astogenetic stage". (Boardman, Cheetham, and Cook, 1970, p. 9). The authors in the discussion following described ways in which dimorphism may be expressed, and these are summarized below:

1) The possession or lack of a given structure.
2) Particular location within a given budding pattern.
3) Difference in size.
4) More complex differences in terms of both structure and function.
Consistent differences between zooecia were not recognized in external or internal examination of type and nontype specimens of H. cryptopora, the type species. There was some variation in the size of zooecial apertures; but no consistent arrangement of larger or smaller polymorphs in the sense of Borg was recognized. A diagram of the frequency distribution of zooecial void diameters was constructed from measurements of randomly selected zooecia in the lectotype and in a single nontype specimen (Text fig. 14A). The diagram shows a continuous and nearly normal distribution with moderately negative skewness.

Gregory (1896, p. 201; 1909, p. 185) referred to the small dimorphs as mesopores and stated that Heteropora was "most closely allied to the genus Heterotrypa", a Paleozoic trepostome. Internal characters as outlined here, however, make as close a relationship to the Trepostomata as suggested by Gregory appear highly improbable. Borg (1933, p. 283) apparently based his diagnosis of

Heteropora on Recent species which he assigned to the genus, in part characterizing the genus as "autozoids . . . not forming clusters; kenozoids smaller and much more numerous than the autozoids, located between them and thus separating them."

The consistent citation of dimorphism as a diagnostic character apparently stems, in most instances, from three factors:

1) Reference to the original figures of Goldfuss which were somewhat idealized, especially magnified views.
2) Reference to Blainville's definition which, again, was derived by examination of figures rather than specimens.
3) Consideration of the characters of species other than the type species.
Remarks on wall structure. - The microstructure in all specimens sectioned was poorly preserved. In thin section, the zooecial walls are light grey to light brown in color, and homogeneous to subgranular with a scattering of small dark grains. Often the boundary between the zooecial wall and calcite infillings of the zooecial chambers was poorly defined (Pl. 33, fig. 2a). Orally convex lineations were, however, sometimes observed (Pl. 35, figs. 1a, $2 \mathrm{a})$. These are interpreted as remnants of originally laminated microstructure.

Light-colored, homogeneous calcite forms irregular bodies in the cortex. This material is probably secondary in origin because it cuts across laminate microstructure unconformably (Pl. 35, fig. 2a). The original zooecial wall tissue is inferred to have been laminate.

Heteropora cryptopora (Goldfuss), 1826
Pl. 32, figs. 1a-f;
Pl. 33, figs. 1, 2a-c; Pl. 34, figs. 1a, b, 2a, b; Pl. 35, figs. 1a-c, 2a, b
1826. Ceriopora cryptopora Goldfuss, Petrefacta Germaniae, vol. 1, p. 33, pl. 10, figs. $3 \mathrm{a}-\mathrm{d}$.
1830. Heteropora cryptopora (Goldfuss), Blainville, Zoophytes: Dictionnaire de Science Naturelles, vol. 60, p. 382.
1834. Hetcropora cryptopora (Goldfuss), Blainville, Manuel d'Actinologie ou de Zoophytologie, p. 417, pl. 70, fig. 4.
1846. Heteropora cryptopora (Goldfuss), Michelin, Iconographie Zoophytologique, description par localites et Terrains des Polypiers Fossiles de France et pays environnants, p. 3.
1851. Ceriopora cryptopora Goldfuss, Von Hagenow, Die Bryozoen der Maastrichter Kreidebildung, p. 53, pl. 5, fig. 6.
1933. Non Heteropora cryptopora (Goldfuss), Borg, Zool. Bidrag från Uppsala, vol. 14, p. 283.
1953. Hetcropora cryptopora (Goldfuss), Voigt, Geol. Staatsiņst. Hamburg, Mitt., vol. 22, pp. 62-3, text fig. 1, pl. 6, fig. 5.

Type. - UB 118a is here designated as the lectotype. UB 118a was figured by Goldfuss, 1826, pl. 10, fig. 3a; Von Hagenow, 1851, pl. 5, fig. 6; Canu and Bassler, 1920, text fig. 222A, p. 681; and here, Pl. 32, figs. 1a-f; Pl. 33, fig. 1.

Type locality and horizon. - Goldfuss (1826, p. 33) cited the locality as "Petersberge bei Maastricht". Rocks exposed there are Upper Cretaceous, Maastrichtian in age.

Material studied. - Five syntypes were borrowed from the Institut für Palaeontologie, Universität Bonn, Bonn. Six thin-sections and six acetate peel replicas on two slides were made from the lectotype, UB 118a. Three thin-sections and three acetate peel replicas were made from one paralectotype numbered collectively with four other paralectotypes as UB 118b. The paralectotype sectioned was figured by Goldfuss, 1826, pl. 10, fig. 3b; and here, Pl. 33, figs. 2a-c, Pl. 34, fig. 1b. Most of both specimens remain as remnants after thin-sectioning.

Thin-sections and acetate peel replicas were also made from 11 specimens collected from the Maastrichtian, Maastricht, Netherlands (USNM Loc. 2387).

Description. -
Growth habit - Zoaria are commonly robust masses with bulbous outgrowths or subcylindrical branches (Pl. 32, fig. 1a). Overgrowth units are lenticular and commonly have moderately to strongly convex distal and concave proximal surfaces (Pl. 32, fig. 1f; Pl. 33, fig. 2b; Pl. 34, fig. 2b). Branches are formed by the synchronous expansion of intrazoarial overgrowths from independent loci (Pl. 33, fig. 2b). The chambers of a few zooecia (commonly located near major growth axis) are continuous from subjacent to suprajacent overgrowth (Pl. 33, figs. 2b, c; Pl. 34, fig. 2b), indicating a boundary between growth phases. The zooecial walls of continuously growing zooecia commonly show thin dark zones followed by thin-walled and often offset growth (Pl. 34, fig. 2b). Laterally, the subjacent zoarial surface is draped by a thin (about .005 mm ), dark-colored basal layer which generally sags partly into the subjacent zooecial chambers ( Pl .34 , fig. 1a). The basal layer is distinctly laminate with laminae directed about parallel to the surface of the basal layer.

Endozone - Zooecia have thin (about .01 mm ), parallel-sided walls (Pl. 34, fig. 1a). The walls are homogeneous to subgranular with thin zooecial linings. Interzooidal pores were rarely observed.

Exozone - Zooecia are nearly straight, gently curved, or slightly undulatory in growth. Zooecial walls are symmetrical in thickness across the zooecial boundary zone. The walls are nearly parallelsided, and show a slight but regular increase in thickness orally. The walls thin near the aperture, and commonly have rounded to acutely lanceolate profiles (Pl. 33, fig. 2a; Pl. 34, fig. 1a; Pl. 35, figs. 1b, 2a). Less commonly, slight variation in wall thickness occurs producing submoniliform cross sections. Interzooidal pores are small in diameter (about .002 to .003 mm ) and are seen infrequently (Text-fig. 14D). Zooecial chambers have elliptical to subelliptical cross sections.

Diaphragms - Intermediate diaphragms are common in zooecia subjacent to overgrowths. The diaphragms are generally seen about 0.1 to 0.3 mm aboral to the aperture ( Pl .32 , figs. $1 \mathrm{f}, \mathrm{g}$; Pl. 33, figs. 2a-c; Pl. 34, figs. 1a, b, 2b). Less commonly, diaphragms are scattered throughout the overgrowth unit (Pl. 34, fig. 2b). The diaphragms generally are thin (about .003 to .006 mm ) and nearly planar, and show slight aboral flexure at the juncture with the zooecial wall before merging continuously with the zooecial lining.

Basal diaphragms were seen rarely, occurring just subjacent to the boundary of zoarial growth phases in zooecia whose chambers are uninterrupted by basal layers.

Brood chambers - Brood chambers occur commonly. The chambers are wide but shallow (see Table 19) and lenticular with slightly convex proximal and distal surfaces. The chambers occur in the more proximal portions of a single overgrowth unit (Pl. 32, figs. 1e, f; Pl. 34, fig. 2b ). Most subjacent zooecia are closed by the brood chamber floor; a few pass through the chamber to the roof. These zooecia are commonly thin-walled and continuous with thin-walled, septate partitions ( Pl .33 , fig. 1). The partitions radiate laterally from the open central portion of the chamber. The brood chamber roof is thick (about .03 to .04 mm ); pores are about .01 mm in diameter.

## TABLE 19

STATISTICAL SUMMARY OF MEASUREMENTS OF HETEROPORA CRYPTOPORA (GOLDFUSS)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | N Zr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zooecial-Exozone } 1 \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 11 | . 03 | . 08 | . 02 | 24 | 75 | 75 | 2 | 1 |
| Zch-CsSn-NMxDn | . 10 | . 02 | . 06 | . 02 | 25 | 75 | 75 | 2 | 1 |
| ZcCh-CsSn-MxDn | 1.8 | 1.0 | 1.2 | . 2 | 13 | 75 | 75 | 2 | 1 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| CdZcWl-Th | . 070 | . 015 | . 034 | . 011 | 32 | 75 | 75 | 2 | 1 |
| ZdPr-Cn/ZcCssn | 2. | 0. |  |  |  | 50 | 50 |  | 3 |
| $\begin{aligned} & \text { Zooecial-Exozone } 2 \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 14 | . 04 | . 08 | . 02 | 26 | 50 | 50 | 1 | 2 |
| ZcCh-CsSn-NMxDn | . 10 | . 03 | . 06 | . 02 | 30 | 50 | 50 | 1 | 2 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{Zc} \mathrm{Ch}-\mathrm{CsSn}-\mathrm{NM} \times \mathrm{Dn}}$ | 2.1 | 1.0 | 1.3 | . 3 | 21 | 50 | 50 | 1 | 2 |

*In millimeters
Exozone 1: Outer exozone, measured in tangential section. Exozone 2: Inner exozone, measured in transverse section.
STATISTICAL SUMMARY OF MEASUREMENTS OF HETEROPORA CRYPTOPORA (GOLDFUSS)


TABLE 20

## FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN HETEROPORA CRYPTOPORA (GOLDFUSS)

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Outer Exozone <br> Regular | 4 | 35 | 9 |  |  |  |  |
| Sub | 7 | 16 |  |  | 1 | 2 |  |
| Irregular |  | 1 |  |  |  |  |  |
| Inner Exozone <br> Regular | 1 | 26 | 3 |  |  |  |  |
| Sub | 9 | 10 |  |  | 1 |  |  |

Irregular

Discussion. - Von Hagenow (1851, p. 53) returned the species H. cryptopora to Ceriopora and singled out only one specimen of the syntype suite to bear the name C. cryptopora. This restriction was made in his synonymy of C. cryptopora by the citation, "Ceriopora cryptopora Goldfuss. Th., Petr. I, p. 33, Taf. X, fig. 3, a (nicht b-d).", and his statement in the remarks following that: "Mir ist nur dieses eine Exemplar in Bonnenser Museum, von Maastricht bekannt." Von Hagenow then assigned the remaining figured specimens (Goldfuss, 1826, pl. 10, figs. b, c, d) to other species of Heteropora.

Gregory (1909, pp. 188-9) carefully reviewed Von Hagenow's analysis of the specimens from the syntype suite of C. cryptopora and remarked that it was "simplest to restore Goldfuss' conception of this species". He considered that all of the specimens of the syntype suite illustrated by Goldfuss (pl. 10, figs. 3a-d) as C. cryptopora were conspecific; and in doing so, he consistently referred to the specimen illustrated by Goldfuss (pl. 10, fig. 3a) as the type.

Gregory's arguments were, in turn, followed by Borg, 1933 (p. 283). Borg stated that:

[^4]

Text-figure 14 A-D. Histograms and cumulative curve from the lectotype and one topotype of Heteropora cryptopora (Goldfuss). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.

Designation of the lectotype here follows the previous restriction of Von Hagenow (1851) and later followed by Gregory (1909) and Borg (1933).

Remarks on morphology. - Diaphragms were interpreted to be intermediate, rather than basal, because a few show slight aboral flexure at the juncture with the zooecial wall.

There is apparently a relationship between diaphragm emplacement and zoarial growth cycles. Intermediate diaphragms commonly occur singly in a zooecium and are commonly emplaced a short distance proximal to the terminal surface of an overgrowth unit. Generally, only zooecia with chambers continuous orally into the superjacent growth unit lack diaphragms.

The taxonomic and phylogenetic significance of the zoarial mode of growth exhibited by $H$. cryptopora is at present unknown. Description and illustrations of a number of species presently thought to be assignable to many diverse genera and families suggests that repetitious addition of intrazoarial overgrowths may be a relatively widespread mode of zoarial growth in cyclostome bryozoans, and some trepostomes as well (Boardman, 1960, pp. 39-40, 57-58, pl. 7, fig. 4, for Leptotrypella multitecta Boardman, a Devonian trepostome). Below is a partial listing of a few species and illustrations suggesting the occurrence of this mode of growth in Cyclostome Bryozoa:

In Canu and Bassler, 1926 - Ceriopora falax, text-fig. 13, p. 28; Diplocava globosa, text-fig. 38, p. 74; Multicrescis lamellosa, textfig. 2, p. 14; Multigalea canui, text-fig. 31, p. 62.

In Gregory, 1909 - Multicrescis tuberosa, text-fig. 54, p. 207; Radiopora neocomiensis, text-fig. 74, p. 285; Reptomulticava fungiformis, text-fig. 39, p. 136.

In this study, incremental addition of intrazoarial overgrowths was seen to play a major role in the zoarial growth of Diplocava incondita and, to a lesser extent, Ceriopora micropora.

Genus LEIOSOECIA Canu and Bassler, 1920
Type species: Multicrescis parvicella Gabb and Horn, 1861, by original designation and monotypy, Canu and Bassler (1920, p. 823).

[^5]1953. Pars Leiosoecia Canu and Bassler, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G72.
Tentative diagnosis. - Zoaria branched; branches with welldifferentiated coaxial exozones and endozones. Distally growing branches commonly intersecting and anastomosing. Zooecia dimorphic.

In exozone, zooecial walls distinctly laminate. In profile section, laminae wavy, sometimes crenulate, forming irregular V - to U shapes convex orally. Zooecial lining present.

Intermediate diaphragms occur occasionally in outer exozone.
Taxa included. - Only the type species, L. parvicella, is here included. Internal characters of other species assigned to Leiosoecia are unknown to me.

Discussion. - Canu and Bassler (1920, p. 823; 1922, p. 99) based their definition on secondary specimens of the type species. These secondary specimens are not considered to be congeneric with the lectotype. The original definition by Canu and Bassler (1922, p. 99) was extremely brief: "Zoarium consisting of cylindrical tubes and regular, parietal mesopores", and could apply to many tubular bryozoans. In 1953, Bassler added the description of the brood chamber to the definition and figured it. This addendum was also based on secondary specimens identified with the type species by Bassler but not considered here to be congeneric with Leiosoecia.

Leiosoecia, in its simplicity of appearance and mode of growth, shows close morphologic similarity to Tetrocycloecia Canu. The morphologic differences between them are considerably less than differences between the other genera studied. In Leiosoecia, however, dimorphism is poorly expressed in zooecial dimensions, and intermediate diaphragms are occasionally seen.

Leiosoecia parvicella (Gabb and Horn), 1861
Pl. 36, figs. 1a-h
1861. Multicrescis parvicella Gabb, and Horn, Acad. Nat. Sci. Philadelphia, Proc., vol. 12, p. 367.
1861. Multicrescis parvicella Gabb and Horn, Gabb, Acad. Nạt. Sci. Philadelnhia, Jour., ser. 2, vol. 4, p. 401, pl. 69, figs. 36-38.
1861. Multicrescis parvicella Gabb and Horn, Gabb, and Horn, Acad. Nat. Sci. Philadelphia, Jour., ser. 2, vol. 5, p. 178, pl. 21, fig. 70.
1905. Multicrescis parvicella Gabb and Horn, Johnson, Acad. Nat. Sci. Philadelphia, Proc., vol. 57, p. 5.
1907. Non Heteropora parvicella (Gabb and Horn), Ulrich, and Bassler, Geol. Sur. New Jersey, Paleont. Ser., vol. 4, p. 327, pl. 23, figs. 1-2.
1920. Non Leiosoecia parvicella (Gabb and Horn), Canu, and Bassler, U. S. Nat. Mus., Bull. 106, p. 823, text figs. 273A-F.
1922. Non Leiosoecia parvicella (Gabb and Horn), Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 100.

Type. - ANSP 31261 is designed as the lectotype. The lectotype is the only syntype known to be preserved. The specimen, ANSP 31261, compares favorably in zoarial form and surface appearance to Gabb and Horn's illustration (Gabb and Horn, 1861, pl. 69, figs. 36-38) and is apparently the specimen referred to by Johnson (1905) as the figured specimen. ANSP 31261 is labeled "Type".

Type locality and horizon. - The label with the lectotype bears the inscription, "Timber Creek, N.J.", one of the two localities originally listed by Gabb and Horn. The specimen probably comes from the Vincentown Formation (pers. comm. H. Richards, 1967) of Paleocene age.

The age and distribution data cited for L. parvicella is here considered questionable. E. O. Ulrich, R. S. Bassler, and others gathered extensive collections of bryozoans from a number of localities in New Jersey and Delaware, including Timber Creek, the cited type locality. These collections, now housed by the National Museum of Natural History and the United States Geological Survey in Washington, D.C., were searched for specimens bearing resemblance to the lectotype; none was found.

Material studied. - The lectotype was kindly loaned to the author by Horace Richards, Academy of Natural Sciences of Philadelphia. Most of the original zoarial fragment remained intact after six thin sections and four acetate peels were made. Duplicate peels are preserved in the National Museum of Natural History collection and the author's collection.

Description of the lectotype. - This description is based solely on the lectotype, the only specimen of $L$. parvicella known. This description includes assessment of nongenetic variation within this colony. No assessment of genetic or other interzoarial variation with L. parvicella is implied in this description.

Growth habit - Branches are roughly cylindrical. The zone of zooecial bending is relatively broad and begins commonly deep in the endozone ( Pl .36 , fig. 1e). Zooecial growth axes are moderately undulatory in both endozone and exozone ( Pl .36 , figs. 1d, e).

Endozone - The zooecial walls are thin and commonly parallel-
sided, sometimes with a small variation in thickness giving submoniliform cross sections (Pl. 36, fig. 1e). Zooecial chambers are polygonal to subpolygonal in cross section. Interzooidal pores are rare. Zooecial walls are homogeneous to subgranular with thin, dark zooecial linings.

Exozone - Large dimorphs are commonly surrounded by small dimorphs, less commonly are directly adjacent to another large dimorph (Pl. 36, fig. 1f).

Zooecial walls of large and small polymorphs are similar in appearance. Both commonly show moderate asymmetry in thickness across the zooecial boundary zone, and both have moniliform profiles due, in part, to the flare of interzooidal pores and to the longitudinal variation in wall thickness (Pl. 36, fig. 1 g ). Walls of large polymorphs, however, generally appear to be more parallelsided in profile. Interzooidal pores were rarely observed (Text-fig. 15C).

Diaphragms - Intermediate diaphragms were observed approximately .02 to .05 mm aboral to aperture in both large and small dimorphs (Pl. 36, fig. 1g). Diaphragms are nearly planar and flex aborally at the juncture with the zooecial wall to merge with the zooecial lining.

TABLE 22
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN THE LECTOTYPE OF LEIOSOECIA PARVICELLA (GABB AND HORN)

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg. |
| :--- | ---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Exozone | 7 | $4+$ | 4 |  | 1 |  |  |
| Regular | 7 | 26 | 3 |  |  |  |  |
| Sub | 2 | 4 |  |  |  |  | 2 |
| Irregular |  |  |  |  |  |  |  |
| Endozone | 2 | 1 | 7 |  |  |  |  |
| Regular |  |  |  | 11 | 2 |  |  |
| Sub |  |  | 2 |  |  |  |  |
| Irregular |  |  |  |  |  |  |  |

TABLE 21
STATISTICAL SUMMARY OF MEASUREMENTS OF LEIOSOECIA PARVICELLA (GABB AND HORN)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | $\mathrm{N} \mathrm{Zr}_{r}$ | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zoarial |  |  |  |  |  |  |  |  |  |
| $\mathrm{Br}-\mathrm{CsSn}-\mathrm{MxDn}$ | 2.8 | 2.6 |  |  |  | 4 |  | 1 | 1 |
| Zooecial-Exozone |  |  |  |  |  |  |  |  |  |
| ZcCh-CsSn-MxDn | . 12 | . 03 | . 08 | . 02 | 23 | 53 | 53 | 1 | 1 |
| ZcCh-CsSn-NMxDn | . 10 | . 02 | . 06 | . 02 | 26 | 53 | 53 | 1 | 1 |
| ZcCh-CsSn-MxDn | 2.1 | 1.0 | 1.4 | . 3 | 19 | 53 | 53 | 1 | 1 |
| Zch-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| CdZcWl-Th | . 10 | . 01 | . 04 | . 02 | 46 | 53 | 53 | 1 | 1 |
| ZdPr-Cn/ZcCsSn . | 1. | 0. |  |  |  | 53 | 53 | 1 | 1 |
| $\begin{aligned} & \text { Zooecial-Exozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 20 | . 07 | . 13 | . 02 | 17 | 46 | 46 | 1 | 1 |
| ZchecsSn-NMxDn | . 14 | . 06 | . 10 | . 02 | 18 | 46 | 46 | 1 | 1 |
| ZcCh-CsSn-MxDn | 1.8 | 1.0 | 1.2 | . 2 | 18 | 46 | 46 | 1 | 1 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |

[^6]STATISTICAL SUMMARY OF MEASUREMENTS OF LEIOSOECIA PARVICELLA (GABB AND HORN)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CdZcWl-Th | . 10 | . 01 | . 04 | . 02 | 42 | 46 | 46 | 1 | 1 |
| ZdPr-Ch/ZcCsSn | 2. | 0. |  |  |  | 45 | 45 | 1 | 1 |
| $\begin{aligned} & \text { Zooecial -Exozone - All Polymorphs } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 20 | . 03 | . 10 | . 03 | 30 | 100 | 100 | 1 | 1 |
| ZcCh-CsSn-NMxDn | . 14 | . 02 | . 08 | . 03 | 35 | 100 | 100 | 1 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NM} \times \mathrm{Dn}}$ | 2.1 | 1.0 | 1.3 | . 2 | 19 | 100 | 100 | 1 | 1 |
| CdZcWl-Th | . 10 | . 01 | . 04 | . 02 | 44 | 100 | 100 | 1 | 1 |
| $\underline{\mathrm{dPr}-\mathrm{Cn} / \mathrm{ZcCsSn}}$ | 2. | 0. |  |  |  | 99 | 99 | 1 | 1 |
| Zooecial - Endozone ZcCh-CsSn-MxDn | . 19 | . 06 | . 13 | . 04 | 27 | 25 | 25 | 1 | 1 |
| ZcCh-CsSn-NMxDn | . 14 | . 04 | . 10 | . 03 | 30 | 25 | 25 | 1 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 1.6 | 1.0 | 1.3 | . 2 | 14 | 25 | 25 | 1 | 1 |
| CdZcWl-Th | . 010 | . 005 | . 007 | . 001 | 20 | 25 | 25 | 1 | 1 |
| ZdPr-Cn/ZcCsSn | 1. | 0. |  |  |  | 25 | 25 | 1 | 1 |
| *In millimeters <br> 1. Lectotype, ANSP 31261. |  | EY TO | CIM | DE |  |  |  |  |  |




Text-figure 15 A-C. Histograms and cumulative curve from the lectotype of Leiosoccia parvicella (Gabb and Horn). A. Normal to maximum crosssectional dimension of a zooecial chamber. B. Ratio of the maximum crosssectional dimension of a zooecial chamber to the normal to maximum crosssectional dimension of a zooecial chamber. C. Count of interzooidal pores per zooecial cross section.

Discussion. - The concept of Leiosoecia, as developed in the descriptions and illustrations of Ulrich and Bassler in Weller (1907), Canu and Bassler $(1920,1922)$ and Bassler (1953), was based on observations of secondary specimens assigned to Leiosoecia parvicella. Thin sections prepared by R. S. Bassler and published illustrations were examined. In addition, a few other secondary specimens from the Paleocene, Vincentown formation, Blackwoodstown, and Vincentown, New Jersey, identified as L. parvicella by Bassler, were thin-sectioned and examined. The secondary specimens (Canu and Bassler, 1920, p. 824, text-figs. 273A-C) are branched, and have coaxial endozones and exozones. The branches, however, do not anastomose, and are small and delicate in appearance in contrast to the robust anastomosing branches of L. parvicella. Zooecia are dimorphic in the secondary specimens (Canu and Bassler, 1920, p. 824, text-figs. 273D-F). The small polymorphs are irregular, nearly sinuous in growth. The large polymorphs commonly exhibit a cylindrical sheath of homogeneous calcareous tissue around the zooecial chamber similar to that seen in Parleiosoecia. There are thick deposits of interzooidal calcareous tissue in the exozone similar in appearance to that commonly seen in hornerids. Intermediate diaphragms are numerous.

Comparison of this preliminary characterization of the secondary specimens to that of the lectotype of L. parvicella reveals significant morphologic differences. These differences are considered to be at least as significant as those which are used to differentiate other genera treated here. Thus, the secondary specimens are not considered here to be congeneric with the lectotype of $L$. parvicella.

Genus PARLEIOSOECIA Canu and Bassler, 1920
Type species: Parleiosoecia jacksonica Canu and Bassler, 1920, by original designation and monotypy.
1920. Parleiosoecia Canu, and Bassler, U.S. Nat. Mus., Bull. 106, p. 824.
1953. Parleiosoecia Canu and Bassler, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G72.
Tentative diagnosis. - Zoaria thin, encrusting expansions, or branched. Branches have strongly differentiated exozones and endozones coaxial with series of hemispherical chambers. Chambers
formed by extensions of basal layer above substrate. Basal layer laminate; laminae inclined proximally oblique. Intersection and anastomosis of distally growing branches common. Zooecia dimorphic.

In endozone, laminate tissue lines zooecial chamber entirely, commonly thickest at proximal tip and deposited directly upon basal layer.

In exozone, cortex of large dimorphs composed of light-colored, homogeneous-appearing calcite.

Walls of small dimorphs distinctly laminate. Laminae commonly broadly arched, orally convex, and continuous across boundaries with adjacent small dimorphs. V-shaped patterns occasionally seen. When adjacent to large polymorphs, laminae abut cortex of large polymorphs at relatively high angle.

Terminal, intermediate, and basal diaphragms present.
Taxa included. - Monotypic for the type species.
Discussion. - P. jacksonica achieves erect branching habit by an unusual modification of the encrusting mode of growth. Locally, the basal layer grows above the encrusted substrate and is extended as a series of hemispherical chambers. The structure produced forms the axial support of a distally growing branch which, in turn, branches to produce a zoarium with ramose growth habit. Branching at short intervals and anastomosis of distally growing branches as seen in zoarial fragments, suggest that the colony so produced was at least moderately large and densely branched in the proximal portions of the colony. Large polymorphs bud from the latero-distal portion of each hemisphere, but apparently never bud from, or overgrow, the distal tip of the branch, i.e., the distal part of the hemi-spheric-axial chamber composed of the basal layer.

The basal layer lacks pores and is structurally different from the calcareous tissue of zooecial walls. Also, the basal layer is often separated from the recumbent zooecial walls by a dark line. The structural characteristics and the size and configuration of the chambers make it difficult to believe that they are zooecial polymorphs.

Similar-appearing axial structures were described by Pergens (1890, p. 318, text-fig. 11) in Cavaria von Hagenow, and in Semilaterotubigera d'Orbigny. Pergens stated that, "leur rôle est inconnu;
peut-être servent-elles à la reproduction" (their function is unknown; perhaps they serve for reproduction). Such a function seems unlikely in Parleiosoecia jacksonica. The only resemblance to brood chambers in other cyclostome species is the formation of a relatively large chamber. The axial chambers differ in position and in structure. Communication pores to adjacent zooecia were not observed, and there are no subjacent zooecia. In P. jacksonica, the wall of the axial chamber is an extension of the basal layer, but brood chamber walls are homologous in structure with zooecial walls and not with basal layers. Brood chamber roofs are commonly (but not invariably) porous. Only a single, central opening was occasionally observed in the axial chambers of $P$. jacksonica. Finally, hollow structures with the typical appearance of brood chambers have been identified in several specimens of $P$. jacksonica; these are located in the exozone and are unrelated to the axial chambers.

The formation of hollow branches by budding of zooecia from a cylindrical-appearing basal layer is known in several cyclostome species such as Seminodicrescis nodosa d'Orbigny, Cavaria ramosa von Hagenow (Gregory, 1899, text-fig. 54, p. 400) and Spiropora macropora d'Orbigny (Pergens, 1890, text-fig. 11, p. 318). Often this may be explained by encrustation on a previously existing structure. In Recent specimens of Densipora corrugata Macgillivray, bits of a tubular, woody, marine plant are often preserved which the bryozoan encrusted. Hamm (1881, p. 25) suggested that axial chambers in Cavaria pustulosa were caused by encrustation on a soft stem. Gregory (1899, p. 399) stated that this conclusion was "untenable", but suggested no alternatives.

The nature of encrustation in P. jacksonica can, at present, be inferred only from negative evidence. Remains that could be interpreted to be a substrate organism or structure have not been recognized in the axial chambers. The outer surface of the basal layer is smooth to rugose, but impressions suggesting encrustation on a substrate organism were not observed. Partitioning of the axial hollow by the basal layer suggests that the cavity was essentially empty when the partition was emplaced. The evidence, as presently understood, suggests that the axial chambers did not encrust a previously existing structure formed by another organism, but that the secreting
epithelium built its own substrate, the basal layer, as growth continued distally.

Parleiosoecia jacksonica Canu and Bassler, 1920 Pl. 37, figs. 1a-g; Pl. 38, figs. 1, 2, 3a, b, 4a-c; Pl. 39, figs. 1, 2a-c, 3, 4, 5; Pl. 40, figs. 1a-f, 2
1920. Parleiosoecia jacksonica Canu and Bassler, U.S. Not. Mus., Bull. 106, pp. 824-5, text-fig. 208e, p. 646; text-fig. 274a-c, p. 825 ; pl. 148, figs. 1-13.
1953. Parleiosoccia jacksonica Canu and Bassler, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, text-fig. 37, figs. 4a, b, p. G72.

Type. - USNM Loc. 2933B-1 is here designated as the lectotype. This specimen is figured here in Plate 37 , figures $1 \mathrm{a}-\mathrm{g}$, and was figured by Canu and Bassler, 1920 (pl. 148, fig. 2).

Type locality and horizon. - The lectotype was probably collected from the Eocene, Jacksonian, Eutaw Springs, South Carolina. The original label with the specimen bears only the word Eutaw.

Material studied. - Five thin-sections and 11 acetate peels were made from the lectotype. Thin-sections and acetate peels were made from two of the paralectotypes: USNM 65447-1 figured by Canu and Bassler in Pl. 148, fig. 1; and here, Pl. 27, figs. 2a-c; USNM 65449 figured by Canu and Bassler in text-fig. 274A, p. 824, pl. 148, fig. 6, and here, Pl. 28, figs. 1a-f.

In addition, thin-sections and acetate peels were made from 31 specimens from Eutaw Springs, S.C., USNM Loc. 2933A. Duplicate acetate peels of all specimens except USNM Loc. 2933A-1-5 are in the author's collection.

There were many inconsistencies between data as given on original labels, as cited in Canu and Bassler (1920), and as cited in catalogue entries for the specimens. Following is a list of the syntype specimens with catalogue numbers and locality data as given on the original labels. Also included is a listing of illustrations believed to be made from a particular specimen, arrived at by direct comparison of the specimen with illustration. The labels on Bassler's thin-sections have plate and figure citations on the original label. It was not possible, however, to identify each section with the cited illustration. In such instances, the citation as originally cited is listed with a question mark.

Cat. No. 65446; 5 thin-sections, Jacksonian, Rich Hill, Georgia.
-1 ? Pl. 148, fig. 9
-2 ? Pl. 148, fig. 10
-3 ? Pl. 148, fig. 11
$-4 \quad$ Pl. 148, fig. 12
-5 Pl. 148, fig. 13
Cat. No. 65447: 2 specimens, Middle Jacksonian, Santee River, 3 miles above Lenuds Ferry, S.C.
-1 Pl. 148, fig. 1
-2 Pl. 148, figs. 4, 5.
Cat. No. 65449: 1 specimen, Eocene ( Jacksonian Middle), 18 miles west of Wrightsville, Johnson Co., Ga., text fig. 274A, p. 824, pl. 148, fig. 6.
No Cat. No.: 4 specimens apparently not catalogued at the same time as the other specimens. Eocene, Middle Jacksonian, Eutaw Springs, S.C.
-1 Pl. 148, fig. 2
Remainder unfigured.

## Description. -

Mode of growth - Branches are commonly cylindrical, occasionally frondose. The distal-most portion of the axial chamber is sometimes open, with the basal layer flexing distally to form a lip around the opening (Pl. 38, fig. 4b; Pl. 39, fig. 4). A narrow zone of zooecial bending separates the well-differentiated exozone and endozone ( Pl . 38, fig. 3a).

Endozone - The walls of large polymorphs are thin and parallel-sided. Large polymorphs are slightly undulatory in their growth, are inclined at a low angle to the branch axis (commonly less than $30^{\circ}$ ), and are long (approximately 1.5 mm from proximal tip to zooecial bend). Large polymorphs but in ranges parallel to a branch axis. Zooecia in adjacent ranges are budded alternately. The overlap of adjacent recumbent zooecia is commonly accomplished by the interwedging of prismatic zooecia which have regular polygonal cross sections (Pl. 39, fig. 5). Sinus and keel accommodation is seen less commonly. The budding pattern is expressed in the exozone by a more-or-less regular, rhombic distribution pattern of large dimorphs (Pl. 39, fig. 2a).

Exozone - Large dimorphs have thin, parallel-sided walls which show little change in thickness throughout the exozone. The walls project slightly above the zoarial surface. Zooecial chambers of
large dimorphs are commonly elliptical to subcircular in cross section. Interzooidal pores are seen rarely.

The longitudinal profiles of the zooecial walls in small polymorphs are somewhat variable in appearance. The walls sometimes show little longitudinal variation in thickness, sometimes show gradual increase orally (Pl. 40, fig. 1f), or show longitudinal, commonly annular, variation in thickness. The walls are commonly thickened subsymmetrically across zooecial boundary zones, but circular or clavate monilar cross sections are often observed. Zooecial chambers have subcircular to elliptical cross sections.

Diaphragms - Terminal diaphragms sometimes occur. Generally, the diaphragms are slightly (about .05 mm ) subapertural in position. The diaphragms have planar oral surfaces and orally convex aboral surfaces. They have short aborally flexed abutments.

Intermediate diaphragms are common in occurrence but frequently show large variability from zoarium to zoarium. Generally, no more than one diaphragm occurs in a single zooecium. Neighboring zooecia commonly have diaphragms in similar positions (Pl. 38, figs. 3a, 3b; Pl. 39, fig. 2b). The diaphragms are thin and planar to slightly convex orally, and they flex aborally at the juncture with the wall to either merge with the zooecial lining, or to form a short, thin abutment distinct from the zooecial lining.

Basal diaphragms were rarely observed, and occur in both large and small dimorphs. The diaphragms are thick, with slightly wavy laminae about parallel to the diaphragm surface. The diaphragms are orally flexed at the juncture with the zooecial wall.

Brood chambers - Brood chambers occur in the middle and outer exozone, and are lenticular to subconical with a planar to domal roof (Pl. 40, figs. 1a-e). The walls of zooecia subjacent to brood chambers often become thin-walled (to about .01 mm ) as much as .1 mm proximally from the brood chamber (Pl. 40, fig. 1d). The brood chamber floor is thin (less than .01 mm ). Most zooecia are sealed at the floor of the brood chamber, but a few large polymorphs pass continuously through the chamber to the roof (Pl. 40, figs. 1b, c, 2). Intra-brood chamber zooecia have thin, parallel-sided walls (Pl. 40, figs. 1c, e). The intra-chamber zooecia are often continuous with vertical septate partitions distributed radially away from the center
TABLE 23
STATISTICAL SUMMARY OF MEASUREMENTS OF PARLEIOSOECIA JACKSONICA CANU AND BASSLER

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF PARLEIOSOECIA JACKSONICA CANU AND BASSLER

| Character | O.R.* |  | X* | $\mathrm{S}^{*}$ | C. V. | N | NZc | NZ r | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| CdZcWl-Th | . 049 | . 012 | . 026 | . 009 | 36 | 20 | 20 | 2 | 5 |
| ZdPr-Cn/ZcCsSn | 0. |  |  |  |  | 20 | 20 | 2 | 5 |
| Zooecial-Small Polymorphs - Exozone ZcCh-CsSn-MxDn | . 09 | . 03 | . 06 | . 01 | 20 | 93 | 93 | 3 | 6 |
| Zch-CsSn-NMxDn | . 06 | . 02 | . 04 | . 01 | 21 | 93 | 93 | 3 | 6 |
| $\underline{\mathrm{ZcCh}}$-CsSn-MxDn | 2.4 | 1.0 | 1.3 | . 2 | 16 | 93 | 93 | 3 | 6 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| ZdPr-Cn/ZcCsSn | 1. | 0. |  |  |  | 93 | 93 | 3 | 6 |
| Brood Chamber BrCh-Wth | 1. |  |  |  |  |  | 1 | 1 | 8 |
| BrChRf-Th | . 038 | . 029 | . 033 | . 004 | 13 | 5 |  | 1 | 8 |
| PrBrChRf-Dr | . 014 | . 001 | . 011 | . 002 | 19 | 12 |  | 1 | 8 |
| *In millimeters $\quad$ KEY TO SPECIMEN CODE |  |  |  |  |  |  |  |  |  |
| KEY TO SPECIMEN CODE |  |  |  |  |  |  |  |  |  |
| 1. Lectotype USNM Loc. $2933 \mathrm{~B}-1$ and 6 topotypes identified by Bassler, USNM Loc. 2933A. <br> 2. Topotypes identified by Bassler, USNM Loc. 2933A-11 (6), -23 (5). |  |  |  |  |  |  |  |  |  |
| 3. Lectotype, USNM Loc. 2933B-1 (50), and topotypes identified by Bassler, USNM Loc. 2933 A <br> 4. Topotypes identified by Bassler, USNM Loc. 2933A-23 (25), -15 (25), -11 (25). |  |  |  |  |  |  |  |  |  |
| 5. Lectotype USNM Loc. 2933B-1 (10), and topotype identified by Bassler, USNM Loc. 2933A-6 |  |  |  |  |  |  |  |  |  |
| 6. Lectotype USNM Loc. 2933B-1 (43), and topotypes identified by Bassler, USNM Loc. 2933A-6 |  |  |  |  |  |  |  |  |  |
| 7. Topotype identified by Bassler, USNM Loc. 2933A-6 (10) |  |  | 8. Paralectotype USNM 65449-1. |  |  |  |  |  |  |



Text-figure 16 A-D. Histograms and cumulative curve from the lectotype and two topotypes of Parleiosoecia jacksonica Canu and Bassler. A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.
of the brood chamber, but leave the central area open. The roof is thick (about .03 mm ) and porous (pores up to .01 mm in diameter). When abandoned, the brood chamber is submerged by a basal layer extending from lateral zooecia, and new zooecia bud from the basal layer.

TABLE 24
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN PARLEIOSOECIA JACKSONICA CANU AND BASSLER

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. | Irreg |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Large Polymorphs - Exozone |  |  |  |  |  |  |  |
| Regular | 3 | 8 |  |  |  |  |  |
| Sub | 7 | 2 |  |  |  |  |  |
| Irregular |  |  |  |  |  |  |  |
| Small Polymorphs - Exozone |  |  |  |  |  |  |  |
| Regular | 2 | 50 | 8 |  |  |  |  |
| Sub | 3 | 20 | 4 |  |  |  |  |
| Irregular |  | 5 |  |  |  |  |  |

Remarks on mode of growth. - Boardman and Utgaard (1966, pp. 1033, 1036, text-fig. 1, p. 1083) reconstructed the three-dimensional shapes of zooecia in several encrusting Paleozoic bryozoans. The recumbent endozonal portions of zooecia commonly developed an interlocking sinus and keel configuration. This configuration was apparently related to the utilization of available space, and was reflected in orderly budding patterns and in the regular arrangement of zooecial apertures as seen at the zoarial surface. Boardman and Utgaard believed that this configuration might be widespread in encrusting tubular bryozoans.

In $P$. jacksonica, the appearance of endozonal zooecia as seen in transverse section, e.g., first row triangular to hemispherical, second row submushroom shape, is generally comparable to the cross sectional shapes of zooecia as described and illustrated by Boardman and Utgaard, suggesting that the recumbent, endozonal portion of zooecia also have an interlocking sinus and keel configuration. In $P$. jacksonica, this pattern is modified because the large polymorphs are oriented radially from, and are recumbent upon, a nearly spherical surface (the axial chamber) rather than a nearly flat surface as seen in the encrusting, sheetlike colonies examined
by Boardman and Utgaard. Accommodation to this spherical surface may, in part, explain the less regular polygonal shapes seen in P. jacksonica (Pl. 39, fig. 5).

Genus REPTONODICAVA d'Orbigny, 1854
Type species: Ceriopora globosa Michelin, 1846, by subsequent designation, Bassler (1935, p. 186).
1846. Pars Ceriopora Goldfuss, Michelin, Iconographie Zoophytologique, Description par Localités et Terrains des Polypiers Fossiles de France et Pays Environnants, p. 246.
1854. Reptonodicava d'Orbigny, Terrain Crétacé Bryozoaires: Paléontologie Française: Description des Animaux Invertébrés, vol. 5, p. 1014.
1896. Pars Ceriopora Goldfuss, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, p. 195.
1909. Pars Ceriocava d'Orbigny, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 127.
1920. Pars Ceriopora Goldfuss, Canu, and Bassler, U.S. Nat. Mus., Bull. 106, p. 678.
1935. Reptonodicava d'Orbigny, Bassler, Fossilium Catalogus, I, Pars 67, Bryozoa, p. 186.
1953. Pars Ceriopora Goldfuss, Bassler, Treatise on Invertebrate Paleontology: Part G, Bryozoa, p. G67.

Tentative diagnosis. - Zoaria globose, hemispherical to cylindrical; sometimes with bulbous to branchlike outgrowths or having less regular, massive shapes. Zoarial surface smooth or uneven, commonly with ridgelike to pustulose monticules. Intrazoarial overgrowths generally covering large areas of growth surface proximally, commonly becoming less common and more localized distally. Endozones are thin and directly adjacent to basal layers or not developed.

In exozone, zooecial walls laminate. Laminae diverging orally at high angle from zooecial boundary zone, arching convex orally across cortex and recurving aborally. Zooecial walls having narrowly integrate appearance in cross section.

Terminal diaphragms and intermediate diaphragms occurring; basal diaphragms common. Intrazooecial spines sometimes seen.

Taxa included. - Only the type species is included here. The internal characters of the second species, R. mamillosa d'Orbigny, assigned to Reptonodicava by d'Orbigny, and species assigned to Reptonodicava by other authors are unknown to me.

Discussion. - Bassler (1935) designated the type species of Reptonodicava without comment. In 1953, Bassler placed Reptono-
dicava in synonymy with Ceriopora following Gregory (1896) and Canu and Bassler (1920). Comparison of thin-sections of the type species of both genera reveals significant differences in morphology. The zooecial wall of $R$. globosa is regular and highly symmetrical but relatively irregular and much less symmetrical in C. micropora Goldfuss, the type species of Ceriopora. R. globosa is characterized by numerous, closely spaced basal diaphragms, but basal diaphragms have not been identified in C. micropora. Intrazoarial overgrowth appears to be a more important means of zoarial increase in $C$. micropora than in $R$. globosa. In C. micropora, the exozonal zooecial walls apparently are composed of orally convex laminae; in $R$. globosa, the walls are probably laminate, but the laminae are recurved from the boundary zone. On the basis of these observable differences in morphology, Reptonodicava and Ceriopora are retained here as separate genera.

Remarks on wall structure. - Microstructure of the zooecial walls was poorly preserved in all specimens available for study. Often, secondary changes have obscured the boundary between the zooecial wall and secondary infilling of the zooecial chamber. Commonly, zooecia appear to have granular or sometimes vaguely laminated structure. Well-defined lineations were observed in a few instances. The lineations, initiated at the boundary zone, are broadly arched and orally convex across the cortex. These lineations are interpreted as remnants of primary lamination. The lamination has been observed to occur only in outer thick-walled portions of the zooecial wall. Presumably, it grades proximally with granular tissue, but its true extent is unknown.

This wall structure is like that seen in Diplocava; also, growth habits show some similarity. Diplocava, however, has numerous peristomial diaphragms and almost no basal diaphragms.

Reptonodicava globosa (Michelin), 1846 Pl. 41, figs. 1a-h; Pl. 42, figs. 1a-g; Pl. 43, figs. 1, 2, 3a-d; Pl. 44, figs. 1, 2a-c
1821. Non Millepora conifera Lamouroux, Exposition Méthodique des Genres de l'Ordre des Polypiers, des Zoophytes d'Ellis et Solander, p. 87, pl. 83, figs. 6, 7.
1824. Non Millepora conifera Lamouroux, Defrance, Dictionnaire de Science Naturelles, vol. 31, p. 84.
1846. Ceriopora globosa Michelin, Iconographie Zoophytologique, Description par Localités et Terrains des Polypiers Fossiles de France et Pays Environnants, p. 246 , pl. 57, fig. 5.
1854. Reptonodicava globosa (Michelin), d’Orbigny, Terrain Crétacé Bryozoaires, Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 1014.
1896. Ceriopora globosa Michelin, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Jurassic Bryozoa, pp. 195-7, text fig. 18, p. 196.
1920. Ceriopora globosa Michelin, Canu, and Bassler, U.S. Nat. Mus., Bull. 106, text-figs. 220A-E, p. 678.

Type. - Sherborn (1940) cited the repository for the specimens described and illustrated by Michelin in the Iconographie Zoologique as the Caen Museum, Caen (Calvados), France. Unfortunately, all of the pre-World War II collections housed at Caen are believed to have been destroyed in the bombardment during the invasion of Normandy. A few of Michelin's specimens have been preserved at the Muséum National d'Histoire Naturelle, Jardin des Plantes, Paris, but Michelin's specimens of Ceriopora globosa are not known to be among them (pers. comm. E. Buge, 1969). Michelin did not designate a holotype, and no specimen is known by me to have been designated as the lectotype.

Type locality and horizon. - Lebissey, Luc, Ranville (Calvados), France; Middle Jurassic, Bathonien.

Material studied. - Specimens from D'Orbigny's collection, identified by D'Orbigny as Reptonodicava globosa, were borrowed from the Muséum National d'Histoire Naturelle, Paris. These specimens were almost certainly studied by D'Orbigny for his concept of the genus Reptonodicava, and it is possible that he had compared these specimens directly with Michelin's primary types of Ceriopora globosa. The specimens (d'Orb. Coll. 2988-1 through 5) display the characters described by Michelin, and resemble his illustrations closely in zoarial shape, surface topography and external appearance of zooecial openings (compare Pl. 42, fig. 1a to pl. 57, figs. 5a, b of Michelin, 1846). In addition, D'Orbigny's specimens were collected at Luc (Calvados), France, one of three localities cited by Michelin. For the above reasons, D'Orbigny's specimens are considered to be conspecific with $R$. globosa in spite of the present impossibility of direct reference to Michelin's primary types.

Thin-sections and acetate peels were made of specimens MNHN d'Orb. Coll. 2988-1 to -3. The card to which the specimens were attached gave the following: Bathonien, Luc (Calvados). Thin-
sections and acetate peels were also made from USNM 32171-1 to -5 from the Bathonien, Ranville (Calvados), and USNM 32180-1 to -4, Bathonien, Langrun. Duplicate acetate peels are preserved in the National Museum of Natural History collection and the author's collection.

Description. -
Mode of growth - Zoaria relatively large (see Table 25). Basal layers of locally occurring intrazoarial overgrowths are relatively thick ( .01 to .02 mm ).

Exozone - Zooecia often grow continuously for long distances (Pl. 43, fig. 3a; Pl. 44, figs. 1, 2a-c). Zooecial walls are symmetrically thickened across zooecial boundary zones. Two major variations are seen locally in the longitudinal profile of the zooecial walls. In one type, zooecial walls show slight longitudinal variation in thickness mainly associated with widely flared interzooidal pores, and resulting in moniliform profiles. Monili are commonly circular, oblong or elliptical (Pl. 43, fig. 3c), less commonly clavate, obovate or sagittate. The monili show little variation in thickness longitudinally resulting in a zooecial chamber with relatively smooth sides overall. When interzooidal pores are uncommon, the zooecial walls locally are parallel-sided (pl. 41, fig. 1e).

On a larger scale, zoaria show repetition of growth zones in which zooecial walls gradually increase in thickness (Pl. 44, figs. 1, $2 \mathrm{a}-\mathrm{c}$ ), producing elongate club-shaped profiles (as modified by local variations described above). Each thick-walled terminal phase is followed by a renewal of thin-walled growth. The boundary zones of major growth phases are commonly marked by concentrations of opaque granules within the wall (Pl. 44, figs. 1, 2a-c). Subzones within each major growth phase are marked by thinner concentrations of opaque granules in the wall (Pl. 44, fig. 2c).

Zooecial chambers commonly have elliptical to subelliptical and smoothly rounded cross sections. Less commonly, short, blunt, intrazooecial spines extend into the chamber producing crenulate outlines. Interzooidal pores are numerous and have large diameters (Table 25, ZdPr-MnDr).

Diaphragms - Terminal diaphragms occur infrequently and are generally seen subjacent to intrazoarial overgrowths, usually with
TABLE 25
STATISTICAL SUMMARY OF MEASUREMENTS OF REPTONODIGAVA GLOBOSA (MICHELIN)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | N Zr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zoarial } \\ & \text { Zr-Wth } \end{aligned}$ | 190 | 80 |  |  |  | 9 |  | 9 | 1 |
| $\begin{aligned} & \text { Zooecial-Exozone } 1 \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 31 | . 09 | . 21 | . 04 | 17 | 75 | 75 | 3 | 2 |
| Zch-CsSn-NMxDn | . 23 | . 07 | . 17 | . 03 | 19 | 75 | 75 | 3 | 2 |
| $\frac{Z \mathrm{cCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{Zc} \mathrm{Ch}-\mathrm{Cs} \mathrm{Sn}-\mathrm{NMxDn}}$ | 2.0 | 1.0 | 1.2 | . 2 | 14 | 75 | 75 | 3 | 2 |
| CdZcWl-Th | . 078 | . 006 | . 033 | . 015 | 46 | 75 | 75 | 3 | 2 |
| ZdPr-Cn/ZcCssn | 4. | 0. |  |  |  | 66 | 66 | 3 | 4 |
| ZdPr-MnDr | . 019 | . 006 | . 012 | . 003 | 27 | 25 |  | 5 | 5 |

[^7]STATISTICAL SUMMARY OF MEASUREMENTS OF REPTONODICAVA GLOBOSA (MICHELIN)

a small space between the diaphragm and the suprajacent basal layer (PI. 43, fig. 3d). The diaphragms are commonly planar, about . 01 to .02 mm thick, and have a short, aborally curved abutment.

Basal diaphragms are thin (about . 001 to .002 mm ), numerous, and often evenly spaced (Pl. 43, fig. 3a; Pl. 44, fig. 2c). The diaphragms may be slightly arched aborally, laminate with two or three laminae, and arched orally at the juncture with the zooecial wall which forms an abutment distinct from the zooecial wall (Pl. 43, fig. 1).

Intermediate diaphragms occur rarely and are seen in the proximal parts of zoaria. The diaphragms are thin (about .001 mm ) and arched orally. They flex aborally at the juncture with the wall to merge with the zooecial lining.

TABLE 26
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN REPTONODICAVA GLOBOSA (MICHELIN)

Circ. Ellip. Ov. Pyr. Polyg. Triang. Irreg.

| Outer Exozone <br> Regular | 5 | 32 | 6 |
| :--- | :---: | :---: | :---: |
| Sub | 6 | 17 |  |
| Irregular | 1 | 15 |  |
| Inner Exozone | 10 |  |  |
| Regular | 4 | 1 |  |
| Sub | 5 |  |  |
| Irregular |  |  |  |

Discussion. - Michelin described Ceriopora globosa as a new species but cited Millepora conifera Defrance (1824, p. 84) as a synonym. This reference was investigated because Ceriopora globosa might be considered as a junior subjective synonym of Millepora conifera.

Defrance (1824, p. 84) had not erected $M$. conifera and clearly referred the name to Lamouroux (1821). Defrance included a restatement of Lamouroux' original description with the addition of his own observations, presumably drawn from specimens which he identified as $M$. conifera Lamouroux.

Perhaps Michelin was referring to specimens which Defrance described as globular varieties of M. conifera; however, Michelin


Text-figure 17 A-D. Histograms and cumulative curve from three topotypes of Reptonodicava globosa (Michelin). A. Normal to maximum crosssectional dimension of a zooecial chamber. B. Ratio of the maximum crosssectional dimension of a zooecial chamber to the normal to maximum crosssectional dimension of a zooecial chamber. C. Compound zoozcial wall thickness D. Count of interzooidal pores per zooecial cross section.
made no statement to that effect. Therefore, only the specimens described by Lamouroux bear on the possibility of synonymy.

Lamouroux' specimens are presumed lost, and a direct comparison cannot be made. In this case, however, Lamouroux' figures are considered to be adequate for comparative purposes. Both the zoarial form and the external appearance of the zooecial openings in $M$. conifera Lamouroux are different from the same characters as illustrated by Michelin for Ceriopora globosa. Thus, Michelin is considered to have made an error in placing Millepora conifera Defrance in synonymy with Ceriopora globosa.

## Genus TETROCYCLOECIA Canu, 1919

Type species: Tetrocycloecia dichotoma Canu, 1919, by original designation, for specimens misidentified by Reuss (1848) as Heteropora dichotoma Goldfuss (1826).
1848. Pars Heteropora Blainville, Reuss, Naturwiss. Abh., vol. 2, p. 35.
1919. Tetrocycloecia Canu, Soc. Géol. France, Bull., ser. 4, vol. 17, p. 346.
1920. Tretocycloecia Canu and Bassler, U.S. Nat. Mus., Bull. 106, p. 826. Obj. svn.
1953. Tretocycloccia Canu and Bassler, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G70. Obj. syn.
1957. Tretocycloccia Canu and Bassler, Buge, Mus. Nat. d'Hist. Nat., Mém., ser. C, Sciences de la Terre, vol. 6, p. 127. Obj. syn.
Tentative diagnosis. - Zoaria branching. Branches have exozones coaxial with endozones, but intergrading through broad zones of zooecial flexure. Zooecia dimorphic.

In outer exozone, zooecial walls indistinctly laminate. Laminae broadly arched, convex orally, and continuous across zooecial boundary zone. Zooecial lining present.

Intrazooecial structures not observed.
Taxa included. - Only the type species, Tetrocycloecia dichotoma Canu, 1919a. The internal characters of other species assigned to Tetrocycloecia are unknown to me.

Discussion. - Canu and Bassler (1920, p. 826) emended Canu's original spelling of Tetrocycloecia to Tretocycloecia. This emendation has been followed by later authors; Buge (1957, p. 127) believed that the emendation was justified because "Tetrocycloecia résultant d'une erreur de transcription du mot grec ayant servi de base au préfixe du nom du genre." Canu, however, used the prefix tetro six times in the original publication:
a) p. 346, original spelling of Tetrocycloeciadae, a new family, repeated twice.
b) p. 346, original spelling of Tetrocycloecia, a new genus repeated in the explanation of pl. 10.
c) p. 346, original spelling of Partetrocycloecia, a new genus.

Thus, there is no evidence in the original publication that could be interpreted as an inadvertent error by Article 32a, ii, of the ICZN, which specifically excludes incorrect transliteration from consideration as an inadvertent error. Evidence for an inadvertent error was not given by Canu and Bassler (1920) when the spelling was emended. The original spelling, Tetrocycloecia, is herein retained; and Tretocycloecia Canu and Bassler (1920) and subsequent authors is considered as an unjustified emendation; therefore, a junior objective synonym of Tetrocycloecia Canu (1919).

Tetrocycloecia dichotoma Canu, 1919 Pl. 45, figs. 1a-g; Pl. 46, figs. 1a-d, 2a-b, 3
1826. Non Ceriopora dichotoma Goldfuss, Petrefacta Germaniae, vol. 1, p. 34, pl. 10, figs. 9a-e.
1848. Heteropora dichotoma (Goldfuss), Reuss, Naturwiss. Abh., vol. 2, p. 35, pl. 5, figs. 20a, b. Mis-I.D.
1877. Heteropora dichotoma (Goldfuss), Manzoni, I Briozoi Fossili del Miocene d'Austria ed Ungheria, Part 3, p. 19, pl. 12, fig. 46. Mis-I.D.
1919. Tetrocycloecia dichotoma Canu, Soc. Géol. France, Bull., ser. 4, vol. 17, p. 346, but non pl. 10, fig. 10.
1920. Non Tretocycloecia dichotoma Canu, and Bassler, U.S. Nat. Mus., Bull. 106, p. 826, text-fig. 275A-I. Mis-I.D., Inv. Emend. Sp.
1920. ? Tretocycloecia dichotoma Canu and Bassler, Canu, Soc. Géol. France, Bull., ser. 4, vol. 19, p. 213. Inv. Emend. Sp., Mis-I.D.?, no text or plate.
1922. Non Tretocycloecia dichotoma Canu and Bassler, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, pp. 108-10, text-fig. 31A, B. Inv. Emend. Sp., Mis-I.D.
1934. Non Tretocycloccia dichotoma Canu and Bassler, Canu, and Lecointre, Soc. Géol. France, Mém., vol. 9, Part 4, pp. 197-8, pl. 38, figs. 1-14. Inv. Emend. Sp., Mis-I.D.
1957. Non Tretocycloecia dichotoma Canu and Bassler, Buge, Mus. Nat. d'Histoire Naturelle, Mém., ser. C, Sciences de la Terre, vol. 6, pp. 127-9. Inv. Emend. Sp., Mis-I.D.
Type. - Six specimens (NMW 1859 L686-1, 2, 3, and NMW 1867xL-1, 2, 3), identified by Reuss as Heteropora dichotoma Goldfuss, are preserved at the Naturhistorisches Museum, Wien.

Specimen NMW 1859 L686-1 bears a close resemblance to Reuss' figure, 1848, pl. 5, fig. 20 (Pl. 45, fig. 1a). The remaining specimens are consistent in external appearance with Reuss' descrip-
tion (1848, p. 35) and figure, and were collected at Eisenstadt, Austria, the type locality. These specimens are not, however, considered here to be syntypic. Reuss commonly listed the date of publication in which a specimen was described, illustrated, or both, as part of the identification number of the specimen (fide A. H. Cheetham). The identification numbers with the specimens studied here do not correspond to any publication of the same date and are considerably later than the original publication in 1848. Although the specimens are not known to be primary types, they are considered to be the most authoritative specimens available. Specimen NMW 1867xL-1 was figured by Manzoni (1877, pl. 12, fig. 46).

Type locality and horizon. - Miocene, Tortonian, Leithakalke; Eisenstadt, Austria.

Material studied. - The specimens, NMW 1859 L686-1, 2, 3, and NMW 1867xL-1, 2, 3, were kindly loaned by Dr. Heinz Kollman, Naturhistorisches Museum, Wien. Three specimens were thinsectioned and peeled:

1859 L686-1. 3 thin-sections, 3 acetate peels. Part of the specimen remained after sectioning.
1859 L686-2. 3 thin-sections, 3 acetate peels. Part of the specimen remained after sectioning.
1867xL1-1. 1 thin-section and 2 acetate peels.
Duplicate peels are preserved in the National Museum of Natural History collection and the author's collection.

Description. -
Mode of growth - Branches are subcylindrical. Zooecia commonly intersect the zoarial surface at $60^{\circ}$ to $80^{\circ}$ ( Pl. 45, fig. 1f; Pl. 46, figs. 1b, 2b).

Endozone - Zooecia are undulatory in growth. Zooecial walls are generally symmetrically to subsymmetrically thickened across zooecial boundary zones, and are parallel-sided to variably thickened with submoniliform to moniliform cross-sections (Pl. 45, fig. 1g; Pl. 46, fig. 2b). Zooecial chambers are commonly subelliptical in cross-section. Interzooidal pores are rare, but are large in diameter (about .01 mm ). Zooecial walls are homogeneous to subgranular in appearance with a thin zooecial lining.

Exozone - Large dimorphs are slightly undulatory and intersect the zoarial surface at $60^{\circ}$ to $80^{\circ}$. The walls of large dimorphs have apertural rims which project slightly above the zoarial surface. Large dimorphs are generally separated by small dimorphs, and commonly are evenly distributed (Pl. 45, figs. 1a-c, e; Pl. 46, figs. 1a, c, 2a, 3). Chambers of large dimorphs have elliptical to subcircular cross sections. Zooecial walls in the inner exozone commonly have subgranular cortices with thin zooecial linings. Lamination becomes more distinct orally, sometimes forming thin zones of laminate tissue which are broadly arched convex orally across the boundary zone. Also, the zooecial lining thickens in the outer exozone to about .02 mm .

$$
\text { TABLE } 28
$$

FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN TETROCYCLOECIA DICHOTOMA CANU*
 zooecial voids were estimated from NMW 1859 L686-1.

Discussion. - Canu (1919) and subsequent authors mistakenly cited the type species of Tetrocycloecia as Heteropora dichotoma Reuss, 1848. Reuss had identified a suite of specimens from Eisenstadt as Heteropora dichotoma (Goldfuss), 1826. Canu (1919, p. 346) stated that "Reuss a confondu cette espèce avec Ceriopora dichotoma Goldfuss 1826", and designated Heteropora dichotoma
STATISTICAL SUMMARY OF MEASUREMENTS OF TETROCYCLOECIA DICHOTOMA CANU

*In millimeters
STATISTICAL SUMMARY OF MEASUREMENTS OF TETROCYCLOECIA DICHOTOMA CANU

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zch-CsSn-NMxDn | . 08 | . 01 | . 04 | . 01 | 26 | 92 | 92 | 2 | 6 |
| ZcCh-CsSn-MxDn | 1.7 | 1.0 | 1.2 | 1.5 | 12 | 92 | 92 | 2 | 6 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| CdZcWl-Th | . 078 | . 019 | . 041 | . 013 | 30 | 92 | 92 | 2 | 6 |
| ZdPr-Cn/ZcCssn | 2. | 0. |  |  |  | 92 | 92 | 2 | 6 |
| $\begin{aligned} & \text { Zooecial-AllPolymorphs - Endozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 17 | . 03 | . 10 | . 03 | 31 | 25 | 25 | 1 | 3 |
| Zch-CsSn-NMxDn | . 11 | . 02 | . 08 | . 02 | 30 | 25 | 25 | 1 | 3 |
| ZcCh-CsSn-MxDn | 2.1 | 1.0 | 1.3 | . 2 | 17 | 25 | 25 | 1 | 3 |
| ZcCh-CsSn-NMxDn |  |  |  |  |  |  |  |  |  |
| CdZcWl-Th | . 026 | . 010 | . 016 | . 004 | 26 | 25 | 25 | 1 | 3 |
| ZdPr-Cn/ZcCsSn | 2. | 0. |  |  |  | 25 | 25 | 1 | 3 |
| $\mathrm{ZdPr}-\mathrm{MnDr}$ | . 008 | . 006 |  |  |  | 3 | 3 | 1 | 3 |

[^8]

ZcCh-CsSn-NMxDn

$\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$
CdZcWI-Th

Text-figure 18 A-D. Histograms and cumulative curve from three topotypes of Tetrocycloecia dichotoma Canu. A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Compound zooecial wall thickness. D. Count of interzooidal pores per zooecial cross section.

Reuss as the type species of Tetrocycloecia. By Article 70B of the ICZN, the primary type specimens of Tetrocycloccia dichotoma were the specimens before Reuss, but by Article 70B, the name assigned to those specimens is Tetrocycloecia dichotoma Canu, 1919.

Gregory (1909, p. 199, footnote 2) believed that Ceriopora dichotoma Goldfuss and T. dichotoma Canu belonged to different genera. This opinion was based on external characters seen in specimens of $C$. dichotoma Goldfuss from the type locality. Canu and Bassler were of the same opinion, and in 1922 (pp. 119-20, textfig. 35) described and illustrated specimens identified as conspecific with $C$. dichotoma Goldfuss from the type locality for which they erected the genus Grammascosoecia. The secondary specimens differ internally in many characters, e.g., bifoliate growth habit, from T. dichotoma Canu as understood here. Until the internal characters of the primary types of $C$. dichotoma Goldfuss have been studied, there must remain some question as to the reliability of existing species concepts as applied to that name. The species concepts, as presently understood, and the temporal separation between $C$. dichotoma Goldfuss and T. dichotoma, however, provide reasonable grounds for continuing to consider $C$. dichotoma and $T$. dichotoma Canu as separate entities.

Canu and Bassler, 1922, p. 108, noted that:
". . . the studies relative to this species have been made from specimens collected in France. We are not entirely certain of our determinations, for we have never been able to procure Austrian specimens for comparison."
A comparison of Reuss' specimens of T. dichotoina to specimens from the Miocene of France figured and identified by Canu and Bassler and later authors, reveals a number of morphological differences (see Canu and Bassler, 1920, text fig. 275A-I; Canu and Bassler, 1922, pp. 108-110, text figs. 31A, B; Canu and Lecointre, 1934, pp. 197-8, pl. 38, figs. 1-14; Buge, 1957, pp. 127-9). The following characters identified in the non-Reuss specimens are not present in the Reuss specimens:

1) Well-defined boundary between exozone and endozone.
2) Zooecia intersect surface at approximately $90^{\circ}$.
3) Endozone walls nonmoniliform.
4) Basal diaphragms in endozonal zooecia.
5) Intermediate diaphragms in exozonal zooecia.
6) Zooecial walls in exozone subsymmetrical to asymmetrical in thickness across the boundary zone.
The specimens from the Miocene of France described by Canu and Bassler (1920, 1922), Canu and Lecointre (1934), and Buge (1957), are here not considered to be conspecific or congeneric with $T$. dichotoma Canu because of the morphotypic differences described above.

## Genus ZONOPORA d'Orbigny, 1849

Type species: Ceriopora spiralis Goldfuss, 1826, p. 36, by original designation and monotypy, D'Orbigny (1849, p. 503).
1849. Zonopora d'Orbigny, Rev. et Mag. Zoologie, vol. 1, ser. 2, p. 503.
1854. Spiroclausa d'Orbigny, Terrain Crétacé Bryozoaires, Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 883. Obj. syn.
1909. Zonopora d'Orbigny, Gregory, Catalogue of Fossil Bryozoa in the Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 2, p. 427.
1922. Spiroclausa d'Orbigny, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 92. Obj. syn.
1953. Zonopora d'Orbigny, Bassler, Treatise on Invertebrate Paleontology, Part G, Bryozoa, p. G71.

Tentative diagnosis. - Zoaria branching; zooecia dimorphic. Large zooecia bud sequentially in helical pattern. Large and small zooecial apertures arrayed in parallel, helical zones. Zone of large zooecial apertures forming a continuous zoarial salient; zone of small zooecial apertures forming a continuous zooecial embayment.

Local intrazoarial overgrowths occur occasionally.
In exozone, outer cortex of zooecial wall light-colored and granular to indistinctly laminate. Laminae broadly curved, convex orally, but commonly not continuous across boundary zone. Zooecial lining thick, dark-colored, with longitudinally directed, undulatory, sometimes crenulate laminae.

Terminal diaphragms and mural spines occurring.
Taxa included. - Only the type species. Internal characters of the types of other species commonly assigned to Zonopora are unknown to me.

Zonopora spiralis (Goldfuss), 1826 Pl. 47, figs. 1a-g; Pl. 48, figs. 1a-g; Pl. 49, figs. 1a-e; Pl. 50, figs. 1a, b, 2a-c, 3
1826. Ceriopora spiralis Goldfuss, Petrefacta Germaniae, vol. 1, p. 36, pl. 11, figs. 2a, b.
1849. Zonopora spiralis (Goldfuss), d'Orbigny, Rev. et Mag. Zoologie, vol. 1, p. 503.
1850. Zonopora spiralis (Goldfuss), d'Orbigny, Prodrome de Paléontologie Stratigraphique Universelle, vol. 2, p. 267.
185+. Spiroclausa spiralis (Goldfuss), d'Orbigny, Terrain Crétacé Bryozoaires, Paléontologie Française Description des Animaux Invertébrés, vol. 5, p. 883, pl. 764, figs. 1-5.
1899. Zonopora spiralis (Goldfuss), Gregory, Catalogue of Fossil Bryozoa in Department of Geology, British Museum (Natural History), The Cretaceous Bryozoa, vol. 1, pp. 427-8.
1922. Spiroclausa spiralis (Goldfuss), Canu, and Bassler, U. S. Nat. Mus., Proc., vol. 61, p. 92-4.

Type. - UB 133 is here designated as the lectotype. The lectotype, labeled "Original zu Goldfuss, 133, Ceriopora spiralis", was firmly cemented to a piece of coquinoid limestone primarily composed of worn bryozoan fragments, echinoid spines and various shell debris. The lectotype is similar in appearance to the specimen figured by Goldfuss, 1826, pl. XI, figs. 2a, b.

Type locality and horizon. - "Petersberge bei Maastricht;" rocks presently exposed here are Upper Cretaccous, Maastrichtian in age.

Material studied. - Dr. A. Durkoop, Geologische Palaeontologische Institut of Bonn, kindly made available the lectotype specimen UB 133. Three thin-sections and four acetate peels on a single acetate slide were made from the lectotype; approximately one-half of the lectotype remained after sectioning.

Four topotypes, collected in the Upper Maastrichtian from Quarry Curfs near Maastricht, were kindly made available for study by Prof. E. Voigt. In addition, 15 topotypes in the collection of the National Museum of Natural History were thin-sectioned and peeled. The specimens were labeled as "Cretaceous, Maastrichtian, Maastricht, Holland, USNM Loc. 2965A".

Duplicate acetate peels of all specimens sectioned are in the collections of the National Museum of Natural History and the author's collection.

Description. -
Mode of growth - Branches have subcircular to elliptical cross sections. Distal to branch bifurcations, the helix pattern of budding
is reversed in one branch. New branches sometimes arise as overgrowths (Pl. 48, fig. 1a).

Endozone - Zooecial walls are thin and parallel-sided. Zooecial chambers commonly have subpolygonal or, less commonly, subelliptical cross sections. Interzooidal pores are rarely seen. Zooecial walls are light-colored and homogeneous to subgranular with thin zooecial linings.

Exozone - Zooecial walls are symmetrical in thickness, or nearly so, across the zooecial boundary zone. Zooecial walls are sometimes nearly parallel-sided (Pl. 50, figs. 1a, b, 3) but more commonly show regular increase in thickness to a maximum which is located slightly suboral to the aperture, producing lancet (Pl. 48, figs. 1c, d, e) or clavate (Pl. 50, fig. 2a) profiles. Locally, most commonly in inner exozone portions of the wall, submonoliform (Pl. 48, fig. 1e; Pl. 49, fig. 1e) and sometimes moniliform cross sections are seen. These result from relatively slight thinning of the wall close to interzooidal pores. Interzooidal pores are narrow (about .01 mm in diameter), straight, and cylindrical for most of their length (Pl. 48, fig. 1g; Pl. 49, fig. 1e). Zooecial chambers commonly have elliptical to subelliptical cross sections.

Mural spines are narrow (about .006 mm in diameter), locally numerous (Pl. 48, fig. 1d; Pl. 49, fig. 1e; Pl. 50, fig. 3), and are most commonly seen in the exozone. The spines have a light-colored core continuous from light-colored tissue in the zooecial wall (Pl. 50, fig. 3 ). In older zooecia, the zooecial lining commonly overlaps and buries the spines ( Pl .49 , fig. 1e).

Diaphragms - Terminal diaphragms are numerous and commonly occur in smaller zooecia. The diaphragms are thin (about . 02 mm ), and pores are about .01 mm in diameter. The walls of zooecia with terminal diaphragms often have constricted apertural rims (Pl. 50, figs. 1a, 2a, b, c). The calcareous tissue of the diaphragms (except where interrupted by pseudopores) is structurally continuous with the zooecial lining of the apertural wall ( Pl .50 , figs. 2b, 2c). The oral surfaces of the diaphragms are flush with the apertures and often appear to form sheetlike deposits when viewed externally (Pl. 47, figs. 1a, b).

## TABLE 29

STATISTICAL SUMMARY OF MEASUREMENTS OF ZONOPORA SPIRALIS (GOLDFUSS)

| Character | O.R.* |  | X* | S* | C. V. | N | NZc | NZr | Spec. Code |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Zoarial } \\ & \text { Br-CsSn-MxDn } \end{aligned}$ | 2.7 | 1.4 | 1.9 | . 4 | 20 | 9 |  | 9 | 2 |
| $\begin{aligned} & \text { Zooecial-Exozone } \\ & \text { ZcCh-CsSn-MxDn } \end{aligned}$ | . 27 | . 04 | . 14 | . 06 | 44 | 36 | 36 | 1 | 1 |
| Zch-Cssn-NMxDn | . 17 | . 03 | . 10 | . 05 | 47 | 36 | 36 | 1 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 2.3 | 1.0 | 1.5 | . 35 | 24 | 36 | 36 | 1 | 1 |
| CdZcWl-Th | . 162 | . 025 | . 072 | . 032 | 44 | 36 | 36 | 1 | 1 |
| $\overline{\mathrm{ZdPr}-\mathrm{Cn} / \mathrm{ZcCsSn}}$ | 4. | 0. |  |  |  | 36 | 36 | 1 | 1 |
| Zooecial - Endozone ZcCh-CsSn-MxDn | . 20 | . 10 | . 15 | . 02 | 17 | 25 | 25 | 1 | 1 |
| Zche-CsSn-NMxDn | . 15 | . 08 | . 11 | . 02 | 14 | 25 | 25 | 1 | 1 |
| $\frac{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{MxDn}}{\mathrm{ZcCh}-\mathrm{CsSn}-\mathrm{NMxDn}}$ | 1.6 | 1.0 | 1.3 | . 16 | 13 | 25 | 25 | 1 | 1 |
| $\overline{\mathrm{ZdPr}-\mathrm{Cn} / \mathrm{ZcCsSn}}$ | 1. | 0. |  |  |  | 25 | 25 | 1 | 1 |
| CdZcWl-Th | . 035 | . 010 | . 018 | . 008 | 44 | 25 | 25 | 1 | 1 |
| ZcWlLn-Th | . 027 | . 002 | . 004 | . 005 | 126 | 25 | 25 | 1 | 1 |

[^9]TABLE 30
FREQUENCY OF VISUALLY ESTIMATED OUTLINES OF ZOOECIAL CHAMBERS IN THE LECTOTYPE OF ZONOPORA SPIRALIS (GOLDFUSS)

|  | Circ. | Ellip. | Ov. | Pyr. | Polyg. | Triang. Irreg. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Exozone <br> Regular |  | 14 | 3 |  | 1 |  |
| Sub | 1 | 12 | 3 |  | 1 |  |
| Irregular |  | 1 |  |  |  |  |
| Endozone Regular | 1 | 1 |  |  | 2 |  |
| Sub | 2 | 8 | 1 |  | 10 |  |
| Irregular |  |  |  |  |  |  |

Remarks on morphology. - The helical growth habit of 7. spiralis poses particular problems of description and interpretation not encountered with other growth habits. One such problem concerns measurement and interpretation of zooecial characters. Certain numerical observations of exozonal zooecia, such as the diameters of the zooecial void, are gathered from tangential sections. The assumptions generally made in the interpretation of such measurements are:

1) If the section is tangential to the surface, all zooecial intersected by the plane of the section will be at approximately the same ontogenetic stage.
2) All zooecia within a given section intersect the section at approximately the same angle (commonly about $90^{\circ}$ ); furthermore, the angle of intersection with the section is about equal to the angle at which zooecia intersect the zoarial surface.
Measurements made from tangential sections of $Z$. spiralis are generally not consistent with either assumption. Measurements, and the statistics generated from those measurements, must be considered biased and only broadly useful in comparison to species satisfying the above assumptions. The bias arises from the helical shape of the branches (refer to Text-fig. 2D). In Z. spiralis, surficial sections parallel to the major axis of branch growth are tangential to the zoarial surface only at the latter-most surface of a zoarial salient (AA' in Text-fig. 20) or, if a deep section is made, are tangential only to the surface at an embayment ( $\mathrm{BB}^{\prime}$ in Text-fig. 20).


Text-figure 19 A-C. Histograms and cumulative curve from the lectotype of Zonopora spiralis (Goldfuss). A. Normal to maximum cross-sectional dimension of a zooecial chamber. B. Ratio of the maximum cross-sectional dimension of a zooecial chamber to the normal to maximum cross-sectional dimension of a zooecial chamber. C. Count of interzooidal pores per zooecial cross section.


A
B


Text-figure 20 A-C. Budding pattern of Zonopora spiralis (Goldfuss). A. An external view of a branch to show the arrangement of large and small zooecia in parallel, helically coiled zooecia. Small zooecia are commonly covered by terminal diaphragms. B. Idealized longitudinal section. Zooecia are curved only in profile through part of any section. In order to show their arrangement, more of the zooecial length is shown in this figure than would be commonly intersected in an actual section. AA' shows the position of a shallow tangential section which intersects only large zooecia. $\mathrm{BB}^{\prime}$ is a deep tangential section large zooecia in deep tangential section, but small zooecia in shallow tangential section. C. Cut away view of a branch. Branch outline indicated by dashed line. Only a single sequentially budded series of large zooecia is shown. Loci of budding indicated by dark helix in central area of branch.

In addition, sections taken along $\mathrm{AA}^{\prime}$ intersect only zooecia with large cross sections; sections taken at $\mathrm{BB}^{\prime}$ intersect thinwalled zooecia with large diameters and clusters of thick-walled zooecia with small diameters. In nearly all sections in which the plane is parallel to the major axis of distal growth, zooecia intersect the plane of the section at a different angle than that at which they intersect the surface. For example, note that zooecia with large cross sections in the middle righthand salient intersect a surface approximately coincidental with $\mathrm{CC}^{\prime}$, rather than at a plane parallel to the major axis of branch growth.

When viewed externally, terminal zooecial coverings sometimes appear to have coalesced as a single calcareous sheet over the zooecial apertures (Pl. 47, figs. 1a, b). If this were so, the calcareous tissue would have been deposited from an outer membrane (gymnocyst of Borg). Thin sections reveal, however, that the diaphragms are typical terminal diaphragms. The diaphragms are not superposed over the zooecial walls but are structurally continuous. The diaphragms were emplaced by tissues aboral to the diaphragm.

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PLATES

## Explanation of Plate 1

Figure Page

1. Ceriocava corymbosa (Lamouroux), 1821 ..... 43MNHN IP2-1, Jurassic, Bathonien; Ranville (Calvados), France.
1a. Branch $\times 1$ with uneven monticular surface; holes areborings.

1 b. Surface of same branch $\times 10$, apparently showing occurrence of large and small polymorphs. Histogram, however, of cross-sectional dimension of zooecial chambers (text-fig. 6A) has unimodal distribution approaching a normal curve.
1 c . Surface of same branch $\times 10$. Note pattern of zooecial openings in upper right monticule and the concentration of smaller zooecia in intermonticular area in center of figure.
1d. Longitudinal thin-section of same branch $\times 10$. Cyclic growth pattern in endozone visible as center of thin, convex zones, caused by annular thickenings of zooecial walls. Note the very thin, undulatory walls and relatively small zooecial diameters in the endozone. Both zooecial walls and zooecial diameters increase in exozone. Tubular structure along branch axis is a boring.
1e. Tangential thin-section of same branch $\times 10$, with normally appearing zooecia in top right and bottom left, and zone of irregular budding in central part of figure.
1f. Transverse thin-section of same branch $\times 10$; all zooecia with terminal diaphragms and thin, closely spaced basal diaphragms. Brood chamber in center of figure; most of endozone removed by a boring organism.
1 g . Longitudinal thin-section $\times 30$. Note circular to oblong moniliform profiles of zooecial wall in inner exozone, and basal diaphragms becoming more numerous and closely spaced in the exozone.
1h. Tangential thin-section $\times 30$. Indistinct laminations concentric with zooecial chamber.


1 b


1 c



1 d
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1 e


## Explanation of Plate 2

Figure ..... Page

1. Ceriocava corymbosa (Lamouroux), 1821 ..... 43MNHN IP2-2, Jurassic, Bathonien; Ranville (Calvados), France.1a. Branch $\times 1$.
1b. Surface of same branch $\times 5$
1c. Surface of same branch $\times 30$.1d. Tangential thin-section $\times 10$; zooecial walls commonlyamalgamate, occasionally integrate.
1 e . Longitudinal thin-section $\times 10$. Thin-walled zooecia with relatively small diameters in endozone. Thick-walled zooecia with large diameter in exozone. Note thick terminal diaphragms and thin, closely spaced, basal diaphragms in exozone.
1f. Transverse thin-section $\times 10$. Illustrates increase in zooecial diameters from endozone to exozone. Note laminate walls in exozone and terminal diaphragms.
1g. Slightly oblique, longitudinal thin-section $\times 10$ with brood chamber in lower left portion of figure. Large cavity to right is a boring.
1h. Longitudinal thin-section $\times 10$. Axillary zone of zooecial intergrowth is shown in upper central part of figure. Zone is just distal to bifurcation of old branch in lower central part of figure. Growth axis of each new branch extends from bottom center to left and right hand corners respectively. Much of endozone in right-hand branch cut by mud-filled boring. Note distal flexure and anastomosis of zooecia growing towards each other from each new branch. Cyclic growth in endozone is illustrated in left-hand branch. Each cycle consists of zooecial walls with long, thin-walled portion capped by annular thick-walled portion; probably represents a single episode of growth.

Explanation of Plate 3


#### Abstract

Figure Page


1-3. Ceriocava corymbosa (Lamouroux), 1821 ......................................
All specimens figured from Jurassic, Bathonien; Ranville (Calvados), France.

1. MNHN IP2-1.

1a. Longitudinal section, acetate peel $\times 50$; zooecial growth direction to the right. Local zone of irregular budding. Zooecial cavities are partitioned by a thick, non-porous diaphragm, unlike numerous basal diaphragms. New zooecial walls are continuous with the oral surface of the diaphragm. Zooecial walls have moniliform profiles, occasional mural spines, and thin basal diaphragms. Calcareous tissue of most of cortex is dark in color; outer cortex is composed of lightcolored tissue outlining zooecial boundary zone. Note numerous basal diaphragms.
1b. Tangential thin-section from same zoarium showing zone of irregular budding.
1c. Longitudinal section, acetate peel $\times 30$. Abandoned brood chamber has floor with interzooidal pores and thick porous roof. Roof is overlain by thin, basal layer and thin-walled zooecia.
2. MNHN IP2-2. Transverse section, acetate peel $\times 30$. Growth direction of zooecia is to the right in zone of irregular budding. Zooecia are thin-walled and irregular in growth direction. Note large open area in central portion of figure.
. 3 USNM Cat. 68941-2, tangential section, acetate peel $\times 30$. Large cavity is interpreted as a brood chamber. Many of the zooecial walls in this figure have an integrate appearance.


1 a


1 b



1 C



## Explanation of Plate 4

Figure ..... Page

1. Ceriocava corymbosa (Lamouroux), 1821 ..... 43MNHN IP2-2, Jurassic, Bathonien; Ranville (Calvados) ${ }_{2}$ France.1a. Tangential section, acetate peel $\times 30$. Zooecial walls general-ly amalgamate in appearance. Many zooecia have smallmural spines.
1b. Longitudinal thin-section $\times 50$; detail of Pl. 2, fig. 1 h, showing anastomosis of orally growing zooecial walls in axillary zones just distal to the bifurcation of a branch. Although zooecial chambers pinch out, zooecial walls merge without break and continue to grow orally.
1c. Transverse thin-section $\times 50$. Laminae generally indistinct, but broad $V$-shaped patterns with apices pointing towards zoarial surface can be seen. Note subapertural position of thick terminal diaphragms.
1d. Longitudinal thin-section $\times 30$. Zooecial walls haye alate to sagittate monilar profiles and are indistinctly laminate. Laminae form $V$-shaped patterns pointing toward zoarial surface. Note correspondence of V -shaped pattern to profile of zooecial wall at skeletal aperture. Basal diaphragms closely and regularly spaced.
1e. Longitudinal section, acetate peel $\times 30$. In endozone, zooecial walls consist of longitudinally repeated two-stage growth cycles: a long, thin-walled growth stage, and a short stage with annular thickenings.

## Explanation of Plate 5


#### Abstract

Figure Page


1-3. Ceriocava corymbosa (Lamouroux), 1821

1. MNHN IP2-1, Jurassic, Bathonien; Ranville (Calvados), France.
1a. Transverse section, acetate peel $\times 100$. Shows thick terminal diaphragms with aborally flexed abutments apparently not merging with zooecial wall. Poorly defined, light-colored patches indicate probable position of pores. Light-colored tissue in outer cortex outlines zooecial boundary zone. Darkcolored tissue, making up most of cortex, is indistinctly laminated. Laminae generally form broad V -shaped patterns with apices pointing towards the zoarial surface.
1 . Transverse section, acetate peel $\times 200$. Shows three thin, intermediate diaphragms just aboral to zooecial aperture. Dark lines sloping aborally from zooecial boundary zone probably represent primary laminate structure. Note granular to nearly structureless calcite along zooecial boundary zone.
1c. Longitudinal section, acetate peel $\times 200$. Zooecial wall in exozone shows indistinct laminate structure. Note lightcolored zone along zooecial boundary zone. Orally flexed basal diaphragm is seen in lower right.
2. USNM 32164-3, Jurassic, Bathonien; St. Aubin (Calvados), France. Longitudinal section, acetate peel $\times 100$. Slightly undulatory, nearly parallel-sided zooecial walls in endozone. Walls are granular. Basal diaphragms flex orally at juncture with zooecial wall and continue orally as zooecial lining for a considerable distance; but commonly, not extending to next diaphragm.
3. MNHN IP2-2, Jurassic, Bathonien; Ranville (Calvados), France. Tangential section, acetate peel $\times 100$. Moniliform profile of zooecial wall shown here results largely from widely flaring profile of interzooidal pores. Section parallels axis of uppermost interzooidal pore, but is slightly oblique or abaxial to other pores which, therefore, appear to be sealed at the zooecial boundary zone. Vaguely spineshaped masses of light-colored tissue project toward the zooecial chamber from the outer cortex, but show little (lower half of figure) or no surficial relief.


1 a





3 a



## Explanation of Plate 6

Figure Page
1-3. Ceriocava corymbosa (Lamouroux), 1821 ..... 43

1. USNM 32164-3, Jurassic, Bathonien; St. Aubin (Calvados), France. Longitudinal section, acetate peel $\times 100$. Laminae have broad, $V$-shaped forms pointing towards the zoarial surface. Light-colored obscurely structured tissue in the outer cortex marks the zooecial boundary zone. Walls are nearly parallel-sided, except where indented by interzooidal pores.
2. MNHN IP2-2, Jurassic, Bathonien; Ranville (Calvados), France. Tangential section, acetate pecl $\times 100$. Zooecial walls show indistinct lamination; thin zooecial linings; short, bluntly rounded zooecial spines; and interzooidal pores with relatively large diameters.
3. MNHN IP2-1, Jurassic, Bathonien; Ranville (Calvados), France.
3a. Longitudinal section, acetate peel $\times 200$. Thick terminal diaphragm is thick and slightly subapertural in position. Basal diaphragms are, by comparison, much thinner. Zooecial walls are indistinctly laminate with laminae orally oblique at a generally higher angle than those in fig. 1.
3 b . Transverse section, acetate peel $\times 200$. The inner cortex is composed of indistinctly wavy to crenulate laminae. Granular to homogeneous calcite in the outer cortex marks the zooecial boundary zone in the inner exozone.
3c. Transverse section, acetate peel $\times 200$. The large chamber is a brood chamber. Subjacent zooecial walls and brood chamber floor are structurally continuous, forming compound wall. The floor is pierced by pores similar in appearance to interzooidal pores. The pores in the roof are sealed proximally by a thin layer of dark-colored tissue which appears to line most of the chamber.

## Explanation of Plate 7

Figure Page

1. Ceriopora micropora Goldfuss, 1826 ............................................... 56

All views are made from the lectotype, UB 119. This specimen was figured by Goldfuss, 1826, pl. 10, figs. 4a, d; von Hagenow, 1851, pl. 5, fig. 13 as Heteropora crassa; and Voight, 1953, pl. 2, fig. 4 as Pennipora beyrichi Hamm.
1a. Zoarial fragment $\times 2$; holes in surface are borings.
1b. Surface of zoarial fragment $\times 30$. Note large variation in size and shape of zooecial apertures. Size variation is continuous, and approaches an unimodal normal curve (Table 7A).
1c. Tangential thin-section $\times 30$. Note large variation in size and shape of cross sections of zooecial chambers. Zooecial walls are broadly amalgamate.
1d. Transverse thin-section $\times 10$ from proximl portion of zoarial fragment. Shows main growth and a single intrazoarial overgrowth.
1e. Longitudinal thin-section $\times 30$; typical appearance of zooecial walls in thin-walled exozone phase. Diaphragm-like structures in oral portions of two zooecia, to right of center, are fractures in the polyester used to imbed the specimen prior to sectioning.
If. Longitudinal thin-section $\times 30$.


1 a


1 c



1 a


1 b


## Explanation of Plate 8

Figure Page

1. Ceriopora micropora Goldfuss, 1826 ..... 56All views are made from the lectotype, UB 119. This specimenwas figured by Goldfuss, 1826, pl. 10, figs. 4a, d; von Hagenow,1851, pl. 5, fig. 13 as Heteropora crassa; and Voight, 1953, pl. 2,fig. 4 as Pennipora beyrichi Hamm.
1a Tangential section, acetate peel $\times 100$. Large, primary interzooecial chamber (brood chamber?).
1b. Longitudinal section, acetate peel $\times 100$. Thin-walled, exozonal growth shows symmetrically thickened, submoniliform to nearly parallel-sided zooecial walls.
1c. Transverse section, acetate peel $\times 100$. Zooecia grow toward the right side of figure. Thin-walled endozonal zooecia are continuous from thick-walled zooecia; intermediate basal layer is not seen here. Some zooecial cavities are continuous.
1d. Thick-walled exozonal growth and intrazoarial overgrowth with new zooecia budding from basal layer. Zooecial walls of subjacent growth show indistinct, broadly curved, orally convex lamination. Endozonal zooecia have granular cortex with thin, laminate zooecial linings.

## Explanation of Plate 9

Figure Page

1. Ceriopora micropora Goldfuss, 1826 ............................................... 56

All views are made from the lectotype, UB 119. This specimen was figured by Goldfuss, 1826, pl. 10, figs. 4a, d; von Hagenow, 1851, pl. 5, fig. 13 as Heteropora crassa; and Voight, 1953, pl. 2, fig. 4 as Pennipora beyrichi Hamm.
1a. Transverse section, acetate peel $\times 200$. Primary laminate structure arches convex orally; laminae appear to be continuous across zooecial boundary zone. Intermediate diaphragm has remnant laminar structure in planar oral portion, but remainder is completely replaced by clear calcite. Note relatively long, aborally flexed abutment which does not merge with zooecial wall.
1 b. Transverse section, acetate peel $\times 200$; primary lamination partly preserved. Large, clear, calcite mass in oral monilis cuts across laminae sharply, and probably results from recrystallization of originally laminar tissue. Also, note intermediate diaphragm in zooecial cavity on right.
1 c. Transverse section, acetate peel $\times 200$. Monili symmetrically to asymmetrically thickened and somewhat variably thickened longitudinally. Primary laminar structure is orally convex. Laminar structure grades aborally into indistinctly structured tissue whic hmay have originally been sublaminate to granular.
1d. Transverse section, acetate peel $\times 100$. Thick-walled exozonal growth shows irregularity in growth direction and variation in wall thickness. Note long tubelike (?) structure in central zooecial cavity. The structure merges with the zooecial wall at oral and aboral ends, and thus appears to be a primary structure.


1 a



 1 b



1 C


2



$3 b$

b

## Explanation of Plate 10

Figure Page
1-3. Corymbopora menardi Michelin, 1846 ..... 64All specimens are from the Cenomanian, Le Mans (Sarthe),France.

1. MNHN Canu Coll. 57057-2 (1).
1a. Distal surface (capitula) of two branches $\times 5$ showinglarge polymorphs.
1b. Lateral view of branch shown above $\times 5$. Note distal ex-pansion forming capitula; small polymorphs are arrangedin rows parallel to the growth axes of the branches.
1c. Detail of 1a $\times 30$ showing shape and arrangement of large dimorphs.
2. MNHN Canu Coll. 57057-2 (2). Surface of stem $\times 30$. Apertures of small polymorphs are arrayed in proximo-distal rows. The walls separating transversely adjacent zooecia form prominent ridges.
3. MNHN Canu Coll. 57057-2 (4).
3a, Transverse thin-section $\times 30$. Shows thin-walled large polymorphs in stem, and thick-walled small polymorphs covering outside of branch completely.
3 b. Longitudinal thin-section $\times 10$ showing stem and expanded capitulum. Chamber in upper right portion of capitulum are lobes of two brood chambers.
3c. Tangential thin-section of stem $\times 30$. Section is slightly oblique. Small polymorphs are seen in the lower part of the figure, and a shallow, longitudinal section of large polymorphs in the upper part of the figure.
3d. Longitudinal thin-section $\times 30$, detail of 3 b . Cross sections of the lobes of two brood chambers can be seen. The positions of two remnant, thick-walled stages are indicated by arrows.
4. MNHN Canu Coll. 57057-2 (3).
4 a . Transverse thin-section of capitulum $\times 30$. Most zooecia exhibit thin-walled phase of growth. The section intersects a thick-walled phase in upper left portion of figure.
4 b. Longitudinal thin-section of stem $\times 30$. Large polymorphs have thin, nearly parallel-sided walls with occasional interzooidal pores. Small polymorphs have short zooecial chambers and thick walls.

## Explanation of Plate 11

Figure Page
1-3. Corymbopora menardi Michelin, 1846 ............................................. 64
All specimens are figured from the Cenomanien, Le Mans (Sarthe), France.

1. MNHN Canu Coll. 57057-2 (5).

1a. Longitudinal thin-section $\times 10$. Cavities in distal portion of capitulum are brood chambers.
1b. Longitudinal thin-section $\times 30$, detail of 1 a . Note prominent interzooidal pores connecting longitudinally adjacent zooecial chamber of small dimorphs, and relatively large interzooidal pores connecting zooecial chamber of small and large dimorphs. Solid arrow is section through wall between transversely adjacent small polymorphs (ridge-forming wall); hollow arrow is section through wall between longitudinally adjacent small polymorphs.
1c. Longitudinal thin-section $\times 30$, detail of 1 a . Shows large polymorph budding from distal wall of small dimorph.
1 e . Longitudinal thin-section $\times 100$, detail of 1 a showing structure of wall between longitudinally adjacent zooecia. Laminae are indistinct but form broadly curved, orally convex patterns in this view.
2. MNHN Canu Coll. 57057-2 (1) $\times 400$. Detail of Pl. 10, fig. 3c. Tangential section of small dimorphs.
3. MNHN Canu Coll. 57057-2 (4).

3 a. Transverse thin-section $\times 100$. Hollow arrow shows position of wall between transversely adjacent small polymorphs (ridge-forming) ; solid arrow shows position of wall between longitudinally adjacent small polymorphs.
3b. Transverse thin-section, crossed nicols $\times 400$. Detail of 3a showing wall structure of small polymorphs. Laminae diverge orally from the longitudinal boundary zone at a low angle in the portion of the wall between transversely adjacent zooecia. (The position of this portion of the wall is indicated by the hollow arrow in fig. 3a). The approximate position of the longitudinal boundary zone is indicated by two hollow triangles. Laminae are slightly arched, convex aborally in this view of the wall between longitudinally adjacent zooecia. (The position of this wall is indicated by the solid arrow in fig. 3a). Compare to the laminar configuration shown in fig. 1c which is at right angles to this view.


$1 b$


1 d


2

$1 e$



3 b


## Explanation of Plate 12

Figure Page
1, 2. Corymbopora menardi Michelin, 1846 ..... 64All specimens are figured from the Cenomanien, Le Mans (Sarthe),France.

1. MNHN Canu Coll. 57057-1. Distal surface of branch (capitulum) $\times 30$ showing brood chamber. Lobes of the brood chamber unite and are continuous with a single zooecial opening in lower portion of the figure.
2. MNHN Canu Coll. 57057-2 (6).
2a. Longitudinal thin-section $\times 10$. Shows two brood chambers and fasciculate appearance of autozooecia in stem and expanded capitulum. Note that expansion takes place as new, large polymorphs bud from the distal wall of small polymorphs.
2b. Tangential thin-section $\times 10$ showing lobate profile of brood chamber.
2c. Detail of $2 \mathrm{~b} \times 100$.
2d. Detail of 2a $\times 100$ showing structure of more distal brood chamber.
2e. Detail of $2 \mathrm{a} \times 100$ showing lobe of more proximal brood chamber.

## Explanation of Plate 13

Figure Page1. Coscinoecia radiata Canu and Lecointre, 193470The lectotype, MNHN Canu Coll. 58872-1, Miocene, Doué-la-Fon-taine (Marne-et-Loire), France, was figured by Canu andLecointre, 1934, pl. 40, figs. 1-4.
1a. Zoarium $\times 1$.
1 b. Surface of zoarium $\times 5$. Large polymorphs are arranged in rows radiating from the monticular areas. Small polymorphs are in monticules and in rows between large polymorphs. The monticule-like structure in the upper right portion of the figure is a brood chamber. Roof with overgrowth is partially broken away revealing hollow interior.
1c. Surface of zoarium $\times 30$. Monticule is shown in upper right. Large elliptical opening in lower left is continuous proximally with a brood chamber.
1d. Longitudinal thin-section of growing tip $\times 10$ showing numerous, closely spaced, basal diaphragms in endozone.
1e. Longitudinal thin-section $\times 10$. Brood chamber in lower portion of figure. Primary growth is encrusted by a cheilostome bryozoan. The cheilostome is, in turn, encrusted by an intrazoarial overgrowth extending from a slightly more distal portion of the primary growth.
1f. Transverse section, acetate peel $\times 5$. Large cavities were made of boring organisms. The primary branch shows encrustation by a cheilostome bryozoan which is, in turn, encrusted from a more distal portion of the original zoarium.
1 g . Longitudinal thin-section $\times 30$. Compare the relatively smooth-sided walls of large polymorphs (at about the center of the figure) to the variably thickened walls of the small polymorphs (at top of the figure). Basal diaphragms occur in zone of zooecial bending and in endozone.
1h. Tangential thin-section $\times 30$. Monticule is shown in right center. Compare pattern of zooecial arrangement to that seen in fig. 1 b . The less distinct pattern in 1 h is probably due to to the difficulty of discriminating between large small-polymorphs and small large-polymorphs in tangential sections.


I a


1 b


1 c

$1 d$

1 e




I c



## Explanation of Plate 14

Figure Page1. Coscinoecia radiafa Canu and Lecointre, 193470
The lectotype, MNHN Canu Coll. 58872-1, Miocene, Doué-la-Fontaine (Marne-et-Loire), France, was figured by Canu and Lecointre, 1934, pl. 40, figs. 1-4.
1a. Longitudinal thin-section $\times 100$, detail of pl. 10, figs. $1 \mathrm{e}, \mathrm{g}$. Shows annularly thickened zooecial walls of small polymorphs with alate monili; also, note broadly curved, orally convex laminae.
1 b. Longitudinal thin-section $\times 100$. The zooecial wall of the large polymorph at lower right shows much less yariation in thickness than the annularly thickened walls of small polymorphs.
1c. Tangential thin-section of intermonticular area $\times 100$. Note vague, pluglike bodies of light-colored tissue.
1d. Tangential thin-section of small polymorphs in monticular area $\times 100$. Note small, blunt, mural spines.
1e. Longitudinal thin-section of zooecial wall shared by small polymorphs $\times 300$. The cortex of most of the wall is composed of light-colored tissue with a dusting of small, dark grains. The structure is obscure, ranging from granular to faintly laminate. Distinctly laminate tissue caps, and partly surrounds, the cortex tissue.
1f. Longitudinal thin-section of large polymorph wall $\times 300$. Wall appearance is similar to 1e, but light-colored cortex tissue here forms discontinuous, pluglike bodies offset to one side of the wall.
1 g . Tangential thin-section $\times 300$. Shows small polymorph with endozooecial spines to left and large polymorph to right. Note pluglike bodies of light-colored, indistinctly laminate cortex tissue bounded by distinctly laminated tissue.
1 h . Tangential thin-section of large polymorph $\times 300$. Calcareous wall tissue is dusted with small, dark granules, and is broadly amalgamate. Note distinct lamination in outer wall, and small pluglike bodies in cortex.

## Explanation of Plate 15

Figure Page

1. Coscinoecia radiata Canu and Lecointre, 1934 ..... 70The lectotype, MNHN Canu Coll. 58872, Miocene, Doué-la-Fon-taine (Marne-et-Loire), France, was figured by Canu andLecointre, 1934, pl. 40, figs. 1-4.
1a. Longitudinal thin-section, detail of pl. 10, fig. 1e, $\times 100$.Zooecia in intrazoarial overgrowth directed proximally in re-lation to major growth axis of branch. Note intermediate dia-phragms and thin basal layer.
1b. Longitudinal thin-section $\times 100$. Shows basal diaphragms in outer endozone, zooecial growth directed towards upper left.
1c. Transverse thin-section $\times 30$. Large cavity is brood chamber; note incomplete floor.
1d. Longitudinal thin-section, detail of $1 \mathrm{f} \times 300$. Shows brood chamber roof and zooecial wall of small polymorph growing to the left.
1e. Tangential thin-section $\times 30$. Large cavity is a brood chamber.
1f. Longitudinal thin-section $\times 3$. Large cavity is a brood chamber.
1g. Longitudinal thin-section of zooecial wall $\times 300$. Calcareous microstructure is laminate except for a small mass of granular tissue in aboral part of cortex.
1 h . Longitudinal thin-section, detail of $1 \mathrm{f} \times 300$. Shows cavity of brood chamber to left, cavity of subjacent zooecia to the right. Lateral growth from oral tips of the subjacent zooecial wall was incomplete. The gap was later sealed by the emplacement of a thick, indistinctly laminate, intermediate diaphragm.
1i. Longitudinal thin-section of a basal diaphragm $\times 300$.


a



1 b


I C


1 g


I h

## Explanation of Plate 16

FigurePage1. Diplocava incondita Canu and Bassler, 192681The lectotype, USNM 69925-2, Cretaceous, Valangian, Ste. Croix(Vaud), Switzerland, was figured by Canu and Bassler, 1926, pl.10, fig. 5 (lower right), and 6.1a. Zoarial fragment $\times 5$. Shows irregular growth habit.
1b. Surface of zoarial fragment $\times 30$. Shows intrazoarial overgrowth on upper right.
1c. Tangential thin-section $\times 30$. Note narrowly integrate appearance of zooecial walls.
1d. Longitudinal thin-section $\times 30$. Much of the primary wall structure altered by diagenetic processes.
1e. Tangential thin-section $\times 30$. Section somewhat deeper than those figured in 1c, f, h. Zooecial walls are amalgamate in this section.
1f. Tangential thin-section $\times 100$. Zooecial walls show both integrate and amalgamate appearance.
1 g . Longitudinal thin-section $\times 100$. Laminae directed orally from boundary zone, broadly recurve aborally and adjoin thin zooecial lining. Apertural rim of zooecial wall is structurally continuous with simple external wall. Also note thin basal diaphragm.
1h. Thin-section transverse to zoarium $\times 30$. Zooecia in center have integrate appearance. Note that most zooecia in overgrowth grow relatively straight from basal layer and are thick-walled.

## Explanation of Plate 17

## Figure Page

1-6. Diplocava incondita Canu and Bassler, 1926
81
All specimens from Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland.

1. USNM Loc. 2404-3, longitudinal thin-section $\times 30$. Shows growth habit with long, continuously growing zooecia uncomplicated by intrazoarial overgrowth.
2. USNM Loc. 2404-1, tangential thin-section $\times 30$. Section passes through a simple external wall in upper right portion of figure.
3. USNM Loc. 2404-8, longitudinal thin-section $\times 30$. Shows mode of growth typified by repetitive overgrowth. Zooecia are initially thick-walled, then slightly thinner, and straight. Two basal diaphragms are seen in third zooecium from bottom.
4. USNM Loc. 2404-5, tangential thin-section $\times 30$. Shallow tangential section in upper right portion of figure shows zooecial walls with integrate appearance. Deep tangential section of intrazoarial overgrowth in lower left portion of figure.
5. USNM Loc. 2404-9.

5a. Longitudinal thin-section $\times 30$.
5 b. Longitudinal thin-section $\times 100$. Shows thick external wall and intrazoarial overgrowth.
5c. Longitudinal thin-section $\times 100$. Zooecial chamber continuous from subjacent growth to overgrowth. Zooecial wall on right apparently grew continuously from subjacent growth to overgrowth, but growth on left was discontinuous.
6. Lectotype 69925-2, figured by Canu and Bassler, 1926, pl. 10. fig. 5 (lower right).

6a. Transverse thin-section $\times 100$. Shows extension of zooecial lining over aboral surface of simple external wall.
6b. Longitudinal thin-section $\times 100$. Zooecial walls show faint recurved lamination. Note thick zooecial linings in zooecia recumbent upon basal layer. Dark line separates two growth episodes.



4


5 c


6 b


## Explanation of Plate 18

Figure Page
1-3. Diplocava incondita Canu and Bassler, 1926 ..... 81All specimens figured from Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland.

1. USNM Loc. 2404-6. Transverse section, acetate peel $\times 200$. Zooecia on left grew orally towards right, have simple external walls, and are encrusted on right by intrazoarial overgrowth. Basal layer and most of wall of overgrowth silicified. Pores through simple external wall in lower zooecium sealed aborally by laminate calcareous tissue continuous with zooecial lining. Note narrow extension of zooecial wall to basal layer of overgrowth. Apertural opening of zooecium in upper left part of figure sealed by basal layer of overgrowth.
2. Lectotype, USNM 69925-2. Longitudinal section, acetate peel $\times 200$. Zooecial wall shows laminae arching orally from boundary zone and recurving aborally. Note apertural tip of zooecial wall extending to basal layer (replaced by silica) of intrazoarial overgrowth. Simple external wall to left separated from zooecial wall by long pore extending obiquely to surface of diaphragm. Pseudopores of simple external wall are sealed aborally by laminate calcareous tissue continuous aborally with zooecial lining and extending orally to form peristome. Also, note thin (one lamina) basal layers in more aboral portions of zooecial chambers.
3. USNM Loc. 2404-7. Tangential section, acetate peel $\times 200$. Zooecial walls have narrowly integrate appearance. Calcareous tissue of peristomial diaphragms merges continuously with zooecial wall. Note laminar structure of peristome (lineations) concentric with restricted apertures in simple external wall.Page
1-3. Diplocava incondita Canu and Bassler, 192681All specimens are figured from Cretaceous, Valangian, Ste. Croix,Switzerland.
4. USNM Loc. 2404-7. Tangential section, acetate peel $\times 200$. Zooecial walls have variously developed integrate appearance. Note deflection of thick zooecial lining at interzooidal pore connecting central and distal zooecial chambers.
5. USNM Loc. 2404-5. Longitudinal section, acetate peel $\times 200$. Zooecia on right grew orally towards left side of figure and have thick, laminate, intermediate diaphragms near aperture. Diaphragms have thick, aborally flexed abutments.
6. USNM Loc. 2404-1. Longitudinal section, acetate peel $\times 500$.
Apertural portion of zooecial wall and oblique section of simple external wall (with circular pores) in lower part of figure overlain by intrazoarial overgrowth. Basal layer and most of proximal zooecial wall of intrazoarial overgrowth is replaced by silica. In subjacent zoarial growth unit, calcareous tissue of zooecial wall is structurally continuous with calcareous tissue of simple external wall (marked by pores). Also, note narrow extension of zooecial wall to basal layer of overgrowth in center of figure marking approximate position of zooecial boundary zone.



I


## Explanation of Plate 20

Figure Page

1. Ditaxia anomalopora (Goldfuss), 1826 ..... 88Lectotype, UB 120, was figured by Goldfuss, 1826, pl. 10, figs.5 c , d; and by von Hagenow, 1851, pl. 4, fig. 9c. Cretaceous,Maastrichtian, Petersberg near Maastricht (Limburg), Nether-lands.
1a. Zoarial surface $\times 30$. Shows monticular areas and irregular distribution of large and small polymorphs in intermonticular areas.
1b. Zoarial fragment $\times 5$. The zoarium is subcylindrical in proximal portion, expanding to frondose distally. Note patchlike distribution of monticules.
1c. Tangential thin-section $\times 30$.
1d. Longitudinal thin-section $\times 30$. Shows bifoliate growth habit; median lamina has dark line at medial boundary zone.
1e. Transverse thin-section $\times 30$.

## Explanation of Plate 21

Figure ..... Page
1,2. Ditaxia anomalopora (Goldfuss), 1826 ..... 88
Cretaceous, Maastrichtian, Guelem, Maastricht (Limburg), Nether-
lands.

1. USNM Loc. 2404-5. Longitudinal section, acetate peel $\times 100$. Shows median layer and recumbent zooecia. Lineations diverging distally from median plane of median layer are interpreted as remnants of primary lamination.
2. USNM Loc. 2404-2.

2a. Longitudinal section, acetate peel $\times 100$. Shows intermediate diaphragms in large polymorph and in small polymorph just distal to it.
2b. Longitudinal section, acetate peel $\times 100$. Large polymorph at top of figure is nearly parallel-sided, but lunarial tissue in proximal wall is obscure because of recrystallization. Proximally, small polymorphs show prominent annular thickenings.
2c. Slightly oblique longitudinal section, acetate peel $\times 50$. Shows intermediate diaphragms in large polymorph at top and center right part of figure. Thick-walled growth in exozone is followed by thin-walled zooecial growth, and a change in zooecial growth orientation suggests a rejuvenated growth phase. Interzooidal spaces, apparently sealed orally and aborally by thin, calcareous walls, are seen between large polymorphs in rejuvenated zone.


1










## Explanation of Plate 22

Figure ..... Page
1, 2. Ditaxia anomalopora (Goldfuss), 1826 ..... 88Cretaceous, Maastrichtian, Guelem, Maastricht (Limburg) , Nether-lands. All photographs taken from acetate peels.

1. USNM Loc. 2405-6. Longitudinal section $\times 100$. Walls of small polymorphs show large increase in thickness from endozone to exozone. In exozone, walls of all small polymorphs in figure show annular thickening at approximately same level.
2. USNM Loc. 2405-5.

2a. Longitudinal section $\times 100$. Large polymorph has thin, nearly parallel-sided walls, although distal walls show some variation in thickness most noticeable just oral to zooecial flexure. Proximal wall of large polymorph has homogeneous calcareous tissue making up a lunaria-like structure.
2 b . Longitudinal section $\times 100$. Large polymorph with thin, nearly smooth-sided zooecial wall and lunaria-like structure in proximal wall. Zooecial walls of small polymorphs are thickened annularly.
2c. Detail of 2a showing wall structure of large polymorph. Proximal wall has homogeneous tissue forming lunaria-like structure. Laminae in outer cortex of adjacent small polymorph dip steeply in aboral direction.

## Explanation of Plate 23

Figure Page

1. Haploecia straminea (Phillips), 1829 ..... 98Lectotype, YM-T81/2, Jurassic, Bajocien (?), Bathonien (?),Yorkshire, England. The thin-sections figured in 1 c -f were madefrom unencrusted branches at less than 5 mm from the growingtip.
1a. Nearly complete zoarium $\times 2$ showing ovoid outline of branching colony.
1 b . Surface of branch $\times 50$. Shows newly-formed simple external walls close to the growing tip. The surfaces of the external walls are slightly depressed relative to the thin apertural parts of the zooecial wall, and zooecia have polygonal, generally hexagonal, outlines.
1c. Longitudinal thin-section $\times 100$. Shows newly-formed simple external wall. Wall is thin and delicate in appearance; compare to more robust appearance of external wall in pl. 24, fig. 1f or pl. 26, figs. 1a, c, d. Zooecial wall shows slight asymmetry in thickness across zcoecial boundary zone and thin zooecial linings. Zooecial wall in right center shows remnant laminae arched orally convex.
1d. Longitudinal thin-section $\times 50$. Shows zone just distal to growing tip where simple external walls first appear.
$1 e$. Tangential thin-section $\times 100$. Shows amalgamate appearance of laminate zooecial walls; also shows interzooidal pores.
1f. Transverse thin-section $\times 50$. Shows central zooecium distinctly larger in diameter than surrounding zooecial openings.


1 c



1 e



1 a


1 b

 1 d


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## Explanation of Plate 24

Figure Page

1. Haploecia straminea (Phillips), 182998Paralectotype YM T81/1, Jurassic, Bajocien (?), Bathonien (?),Yorkshire, England.
1a. Longitudinal thin-section $\times 30$ showing abandoned growing tip and intrazoarial overgrowth. Zooecia in abandoned branch tip are thick-walled and have simple external walls.
1b. Transverse thin-section $\times 30$ showing thick walls in exozone, thin walls in endozone. At top center, peristome protrudes into overgrowth.
1c. Longitudinal thin-section $\times 30$ showing narrow zooecia in endozone with diameters widening distally in exozone. Zooecia of primary branch have simple external walls.
1d. Tangential thin-section $\times 100$. Large zooecium at centerleft shows interzooidal pore to left and more closely-spaced pseudopores in terminal diaphragm to right.
1e. Tangential thin-section $\times 100$ showing simple external walls.
1f. Longitudinal thin-section $\times 100$ showing structure of zooecial wall in exozone and simple external wall. Boundary with thin basal layer of overgrowth is marked by a distinct, dark line.

Explanation of Plate 25


#### Abstract

Figure Page


1, 2. Haploecia straminea (Phillips), 1829
98
Both specimens from Jurassic, Bajocien (?), Bathonien (?), Yorkshire, England. All photographs taken from acetate peels.

1. YM T81/1, paralectotype.

1a. Intersection of two different branches $\times 50$. Curved line from top left to lower right side of figure marks intersection between distally growing branch below (longitudinal section) and side of branch above (transverse section). Complex growth pattern in lower right is an intrazoarial overgrowth of branch to left. Zooecia in branch to upper right had thick simple external walls apparently in place before intersection occurred. Thin, zooecial seals of left-hand branch were apparently deposited at about the same time as intersection occurred (detail in figs. 1c and 1d).
1b. Longitudinal section $\times 200$. Shows basal diaphragms in inner exozone. Only distal-most diaphragm is well preserved. Diaphragms are flexed slightly in oral direction, but appear to abut, rather than merge with, zooecial wall.
1c. Detail of $1 \mathrm{a} \times 200$. Shows boundary between intersecting branches (indicated by arrow). Zooecium on right is sealed by a simple external wall. Thin-walled zooecia on left are sealed by nonporous diaphragms composed of light-colored, granular tissue (directly adjacent to plane of intersection) continuous with cortex of zooecial wall, and lined aborally by laminated tissue continuous with zooecial lining.
1d. Detail of $1 \mathrm{a} \times 200$ showing boundary between intersecting branches (indicated by arrow). Thick-walled zooecium on left is sealed by a thin diaphragm which flexes aborally to merge with zooecial lining. Faint line and thin gap separate both branches. Zooecium on right has a thick, poorly-preserved simple external wall.
2. YM T81/2, lectotype.

2a. Longitudinal section near growing tip $\times 200$. Thin, newlyformed external wall merges continuously with calcareous tissue of zooecial wall.
2b. Longitudinal section near growing tip $\times 200$. Shows thin, newly-formed simple external wall. Calcareous tissue of wall flexes aborally to merge continuously with zooecial wall.





1 a
(14. 1


1 b


## Explanation of Plate 26

Figure Page

1. Haploecia straminea (Phillips), 1829

Paralectotype from Jurassic, Bajocien (?), Bathonien (?), Yorkshire, England. All photographs taken from acetate peels.
1a. Longitudinal section $\times 200$. Shows single zooecium with simple external wall on left, separated by a dark line (indicated by an arrow) from intrazoarial overgrowth in transverse section on far right. Calcareous tissue of simple external wall is light-colored (probably granular) just subjacent to dark boundary line, but distinctly laminate in aboral part; laminae are moderately convex in aboral direction. Basal layer and cortex of overgrowth have lightcolored, homogeneous calcareous tissue. Note thick, distinctly laminate zooecial lining.
1b. Transverse section of branch on bottom $\times 100$. Shows oblique longitudinal sections of intrazoarial overgrowths, each marked by simple external walls. Note basal diaphragm in peristome of lower left-hand zooecium, and intermediate diaphragm abutting simple external wall in zooecium at middle of figure. Poorly preserved lineations of cortex tissue in zooecial walls of bottom branch are interpreted as original laminations directed convex orally.
1c. Longitudinal section $\times 200$. Zooecium on right with simple external wall (peristome directed obliquely towards upper left corner) is separated by two dark lines and a gap from intrazoarial encrustation to left (indicated by arrow). Calcareous tissue of peristome merges continuously with subjacent zooecial wall. The peristome and external wall have a laminate inner portion continuous with zooecial lining, and light-colored, subgranular outer portion continuous with the cortex of the zooecial wall.
1d. Longitudinal section on left and obliquely transverse section of intrazoarial overgrowth on right $\times 200$. Peristomes of zooecia on left are directed almost horizontally to right. External walls are continuous structurally, with calcareous tissue of subjacent zooecial walls. Dark line marks boundary between growth phases.

## Explanation of Plate 27

Figure Page
1-3. Haploecia multilamellosa (Canu and Bassler), 1926 ..... 106All specimens figured were identified by R. S. Bassler and werecollected from the type locality in the Cretaceous, Valangian,Ste. Croix (Vaud), Switzerland.

1. Lectotype, USNM 69922-1, figured by Canu and Bassler, 1926, pl. 9, figs. 1 (center specimen), 5, 6. Most of the internal structure is obscured by recrystallization and silicification.
1a. Branch $\times 5$.
1b. Surface of branch $\times 30$. Zooecia are arranged in rows; zooecial apertures in adjacent rows alternate in position. Peristomes are commonly located centrally with respect to the zooecial walls.
1c. Longitudinal section, acetate peel $\times 30$. Zooecia intersect zoarial surface obliquely; zooecia encrusting primary branch grow orally in approximately the same direction as underlying zooecia.
1d. Tangential section, acetate peel $\times 30$. Zooecial openings are commonly elliptical, with greatest dimension parallel to the growth axis of the branch.
1e. Transverse section, acetate peel $\times 30$. Shows primary branch and single intrazoarial overgrowth. A single, enlarged zooecium occupies the center of the endozone.
2. Paralectotype, USNM 69922-2.
2a. Branch $\times 5$.
2b. Surface of branch $\times 30$. Arrangement of apertures are not as regular as those seen in Pl. 27, fig. 1a.
3. Topotype identified by R. S. Bassler, USNM 2384-1.
3a. Longitudinal section, acetate peel $\times 30$. Shows growing tip of primary branch with a thin, intrazoarial overgrowth.
3b. Tangential section, acetate peel $\times 30$. Tangential to zooecia in primary branch, shows shape and arrangement of zooecial openings.
3c. Transverse section, acetate peel $\times 30$. Shows primary branch with enlarged central zooecium and three generations of intrazoarial overgrowth (upper right). All zooecia, except in outer-most overgrowth, have simple external walls.
3d. Tangential section, acetate peel $\times 30$. Section is tangential to primary branch in axial portion, and longitudinally oblique to intrazoarial overgrowth on either side. Zooecial apertures are arranged in longitudinal rows; zooecia in adjacent rows generally alternate in position longitudinally. Smaller round opening at right center is a peristome.


1 b


2 a


2 b

$1 d$


1 e



1 a

$1 b$


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$1 e$


## Explanation of Plate 28

Figure
1-3. Haploecia multilamellosa (Canu and Bassler), 1926 ..... 106

All specimens figured were identified by R. S. Bassler, and were collected from the type locality in the Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland.

1. USNM Loc. 2384-2.

1a. Longitudinal section, acetate peel $\times 30$. Shows primary branch with coaxial endozone and exozone enveloped by two intrazoarial overgrowths. Zooecium in upper portion of figure has a prominent peristome submerged beneath basal layer of overgrowth.
1b. Tangential section, acetate peel $\times 30$. Section intersects a primary branch (upper right) and an intrazoarial overgrowth. In the lower portion of the figure, two circular peristomes from the subjacent primary branch extend into the overgrowth.
1c. Tangential section, acetate peel $\times 100$. Shows pseudopores and restricted aperture in simple external wall. Section is slightly oblique; lower portion of figure shows thick, exozonal zooecial walls of primary branch; upper part of figure shows thin endozonal walls of intrazoarial overgrowth.
1d. Longitudinal section, acetate peel $\times 100$. Detail of 1 a showing wall structure. Note thin band of light-colored tissue in outer cortex which marks the zooecial boundary zone, and the rodlike extensions from the cortex forming prominent, intrazooecial spines. Zooecia of the primary growth are sealed by simple external walls, and are encrusted by the basal layer of intrazoarial overgrowth. A thin, light-colored boundary zone separates the two (detail, Pl. 31, fig. 2b).
1 e. Tangential section, acetate peel $\times 100$. Section is just aboral to the simple external wall. Note the prominent mural spines extending from the cortex. The spines are partially covered by zooecial lining. Interzooidal pore shown in lower left is narrow, with little flare at either end.
2. USNM Loc. 2384-1. Transverse section, acetate peel $\times 100$. Shows the large central zooecium which lacks zooecial lining. Smaller surrounding zooecia typically have a thin zooecial lining. In exozone, zooecial walls have a few small, mural spines; zooecia are sealed by simple external walls which merge continuously with the zooecial lining.
3. USNM Loc. 2384-10.

3a. Tangential section, acetate peel $\times 100$. Shows typical arrangement of zooecia and shape of openings. Note thick zooecial lining which covers most intrazooecial spines completely. The spines appear as light-colored rods contrasting with the dark-colored, longitudinaly laminated tissue of the zooecial lining.
3 b . Longitudinal section, acetate peel $\times 100$. Shows primary branch and intrazoarial overgrowth along left side of figure. In primary branch, zooecia grow towards upper left and have simple external walls. In the lower left zooecium, the peristome is low, and the peristomial aperture is sealed by calcareous tissue continuous with the simple external wall. The basal layer of the overgrowth is poorly differentiated from subjacent calcareous tissue of the primary branch; but, generally, a thin, light-colored boundary zone separates the two.

1-3. Haploecia multilamellosa (Canu and Bassler), 1926
All specimens were identified by R. S. Bassler and were collected from the type locality in the Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland.

1. USNM Loc. 2384-10.

1a. Longitudinal section, acetate peel $\times 5$. Shows intersection of a distally growing branch with a second branch. The plane of intersection is visible as a dark line (figs. 1c, 1d for detail). Note that the intersection occurred before either primary branch was covered by an intrazoarial overgowth. The overgrowth on the left-hand branch partially envelops the intersecting branch.
1b. Tangential section, acetate peel $\times 30$; at intersection shown in 1a. Plane of intersection appears as a dark line. Overgrowth zooecia are visible in upper right.
1c. Longitudinal section, acetate peel $\times 30$, detail of 1 a. Shows primary branch in center, intrazoarial overgrowth to the left, and intersecting primary branch on right. Zooecial details of intersecting branch are obscure because of silicification of the zooecial walls.
1d. Longitudinal section, acetate peel $\times 30$; detail of 1c. Diaphragms were deposited by zooids on both sides of the zone of intersection. To the left are simple external walls. Diaphragms on the right are thin and non-porous.
1 e . Longitudinal section, acetate peel $\times 100$ showing thin, intermediate diaphragms in intrazoarial overgrowth.
2. USNM Loc. 2384-8.

2a. Longitudinal section, acetate peel $\times 5$ showing oblique intersection of two distally growing branches. In contrast to 1a, the intersection apparently occurred at the actively growing tips of both branches. Note the deflection of endozonal zooecia so that the major axis of distal growth of each branch assumes the approximate orientation of the other branch. Details are shown in 2 b and 2c.
2 b . Detail of 2a $\times 30$. Proximal zooecia are sealed off at the zone of intersection (see 2c). Distally, zooecia merge continuously along the zone of intersection and curve away from the zone of intersection forming a common endozone for a short distance.
2c. Detail of $2 \mathrm{~b} \times 100$ showing closure of zooecia along the zone of intersection. Note the slight distal flexure and thinning of zooecial walls just aboral to the zone of intersection.
3. USNM Loc. 2384-1. Longitudinal section, acetate peel $\times 100$. Shows primary growth on lower left and intrazoarial overgrowth on the upper right. Zooecial wall of the overgrowth have light-colored (interpreted as granular) cortices and thin, dark-colored zooecial linings which merge continuously with the basal layer. The basal layer is difficult to separate from the encrusted zooecial wall and peristomial diaphragm of the primary branch, but a thin, light-colored zone generally marks the boundary.



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## Explanation of Plate 30

Figure ..... Page
1-3. Haploecia multilamellosa (Canu and Bassler), 1926 ..... 106
All specimens figured were identified by R. S. Bassler and were collected from the type locality in the Cretaceous, Valangian, Ste. Croix (Vaud), Switzerland.

1. USNM Loc. 2384-2.
1a. Transverse section, acetate peel $\times 200$. Shows primary branch and two consecutive intrazoarial overgrowths. Overgrowths are one and one-half to two zooecial diameters thick. Intrazoarial overgrowths are separated from each other and from the primary branch by a dark line or by a thin, light-colored boundary zone (indicated by arrows). The simple external wall of the zooecium in the center of the figure is thin with dark laminae similar in thickness and appearance to zooecial lining. The wall is apparently continuous with the zooecial lining to left, but separated on right by a pseudopore. Wall is laminate; basal layer of suprajacent overgrowth is light-colored with indistinct structure interpreted as granular.
1b. Longitudinal section, acetate peel $\times 200$. Shows primary branch on right and two intrazoarial overgrowths, each separated by a thin, light-colored boundary zone (indicated by arrows). Zooecia grew towards upper left. Zooecial apertures are sealed by simple external walls.
2. USNM Loc. 238+-7. Longitudinal section, acetate peel $\times 50$. Shows interruption of zooecial growth in endozone just below center of figure, and renewed growth indicated by offset of zooecial walls. Also note variation in size of central zooecium in lower portion of figure; diameters of other endozonal zooecia show relatively constant diameter.
3. USNM Loc. $2384-10 \times 200$. A thin, intermediate diaphragm is shown in the central zooecium of the intrazoarial overgrowth. The diaphragm shows a slight aboral flexure at juncture with zooecial wall. Primary branch is seen along right side of figure with intrazoarial overgrowth to the left. The primary branch and the basal layer are separated by a thin, light-colored zone (indicated by arrow). Note the small intrazooecial spines in the basal layer. Note slightly undulatory growth and occasional small mural spines typical of endozonal appearance; also, the relatively large interzooidal pore in the zooecial wall at lower center.

## Explanation of Plate 31

FigurePage1, 2. Haploecia multilamellosa (Canu and Bassler), 1926 ..... 106All specimens were identified by R. S. Bassler and collected fromthe type locality in the Cretaceous, Valangian, Ste. Croix (Vaud),Switzerland.

1. USNM Loc. 2384-10.
1a. Transverse section, acetate peel $\times 200$. Shows proximal growth separated from overgrowth by a dark line and, sometimes, by a thin, light-colored boundary zone (indicated by arrow). Simple external wall of zooecium in proximal growth is thin and composed of dark laminate tissue which recurves aborally to merge with zooecial lining, contrasting with lighter-colored and homogeneous calcareous tissue of superposed zooecial walls interpreted here as originally granular. Note thin, dark-colored zooecial linings of recumbent overgrowth zooecia.
1b. Longitudinal section, acetate peel $\times 200$. Shows oral growth of zooecium on right directed toward left-hand side of figure. Zooecium is capped by a simple external wall and encrusted by intrazoarial overgrowth. Zooecia in overgrowth grow orally towards upper left. Laminate tissue of simple external wall flexes aborally to merge with zooecial lining; note oblique section of pseudopore in proximal portion of external wall which appears to separate external wall from compound zooecial wall. Recumbent zooecial walls of overgrowth are very thin, and are separated from proximal growth by a thin, light-colored zone (indicated by arrow).
2. USNM Loc. 2384-2.
2a. Longitudinal section, acetate peel $\times 500$. Shows apertural portion of zooecium on right with simple external wall (middle of figure), and recumbent zooecia of intrazoarial overgrowth on left. Aperture in peristome is sealed by laminate calcareous tissue merging continuously with calcareous tissue of external wall. Recumbent wall of the overgrowth zooecium is separated from simple external wall by thin, light-colored zone (indicated by arrow).
2b. Longitudinal section, acetate peel $\times 500$. Detail of Pl. 28, fig. 1d. Zooecia on left grew orally toward right side of figure, capped by simple external wall (from center top to center bottom of figure) and encrusted by intrazoarial overgrowth. The zooecium in the overgrowth grew orally towards top of figure. The overgrowth is separated from the subjacent zooecial wall and peristomial diaphragm by a thin, light-colored boundary (indicated by arrow). Note the prominent mural spines in the zooecial wall to the left. The spines have cores composed of homogeneous-appearing calcareous tissue, and are generally submerged beneath laminated zooecial lining. A few light-colored cores of small mural spines can be seen in the wall of the recumbent zooecium on the right; the cores are nearly completely submerged beneath the zooecial lining.


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## Explanation of Plate 32

Figure Page

1. Heteropora cryptopora (Goldfuss), 1826 ..... 115Lectotype, UB 118a, figured by Goldfuss, 1826, pl. 10, fig. 3a; byvon Hagenow, 1851, pl. 5, fig 6; and by Canu and Bassler, 1920,text fig. 222A, p. 681. From Cretaceous, Maastrichtian, St.Petersberg near Maastricht (Limburg), Netherlands.
1a. Zoarium $\times 2$. Zoarium is massive with bulbous to digitate branches.
1b. Surface of zoarium $\times 30$. Shows zooecial apertures with nearly continuous variation in size.
1c. Tangential thin-section $\times 30$. Shows zooecial openings which are commonly elliptical to subelliptical, and have nearly continuous variation in size.
1d. Transverse thin-section $\times 30$. A dark line separates zoarial growth phases. Some zooecial chambers subjacent to overgrowth are filled with dark-colored, fine-grained sediment; other zooecial chambers are filled with clear calcite.
1 e . Longitudinal thin-section $\times 10$. Shows zoarial growth by repetitive addition of superposed intrazoarial overgrowths. Chambers in lower right and upper left are brood chambers; both are located proximally with respect to individual growth phases.
1f. Longitudinal thin-section $\times 30$, detail of 1 e . Shows structure of brood chamber. In this view, five zooecia pass through chamber to roof. Intermediate diaphragms are seen in many zooecia suboral to basal layer of overgrowth. Two intermediate diaphragms can also be seen in lower right portion of figure.

1, 2. Hełeropora cryptopora (Goldfuss), 1826
Both specimens figured were collected from the Cretaceous, Maastrichtian, St. Petersberg near Maastricht (Limburg), Netherlands.

1. Lectotype, UB 118a, figured by Goldfuss, 1826, pl. 10, fig. 3a; by von Hagenow, 1851, pl. 5, fig. 6; and by Canu and Bassler, 1920, text fig. 222A, p. 681. Longitudinal thin-section $\times 100$ showing structure of brood chamber. Zooecia and septate structures pass through the chamber. These intrachamber structures have thin, dark-colored walls in contrast to the lighter-colored tissue of the walls distal to the brood chamber. Septate structure on upper left is continuous with intra-chamber zooecium in oblique section. The roof is porous and separated from the superjacent encrusting growth by a thin, dark line.
2. Paralectotype, UB 118b, figured by Goldfuss, pl. 10, fig. 3 b .
2a. Longitudinal thin-section $\times 100$ showing characteristic appearance of zooecial wall. Thin intermediate diaphragms are shown just aboral to apertures of zooecia subjacent to overgrowth phase on right.
2b. Longitudinal thin-section $\times 10$ showing characteristic appearance of zoarial growth by repetitious increments of intrazoarial overgrowths. The outline at the terminal surface of each unit is emphasized by distally convex, dark lines caused by emplacement of intermediate diaphragms roughly coincidental with neighboring zooecia, and by emplacement of a thin, dark basal layer over the subjacent zoarial surface. Branching begins with the development of two separate, but later confluent, growth units on the same zoarial surface.
2c. Longitudinal thin-section $\times 30$, detail of 2 b .



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## Explanation of Plate 34

Figure Page
1, 2. Heteropora cryptopora (Goldfuss), 1826 ..... 115

1. Paralectotype, UB 118b, from the Cretaceous, Maastrichtian, St. Petersberg near Mastricht (Limburg), Netherlands; figured by Goldfuss, 1826, pl. 10, fig. 3 b .
1a. Longitudinal section, acetate peel $\times 100$ showing characteristic profile of apertural parts of zooecial walls. Intermediate diaphragms occur at nearly the same level in all zooecia, slightly aboral to the terminal growth surface. The basal layer of the encrusting growth unit is thin and dark in color. This view of the overgrowth shows the thin-walled, parallel-sided appearance typical of zooecia in the endozone.
1b. Transverse thin-section $\times 30$. Growth phases are separated by two dark lines. The line on the right is caused by intermediate diaphragms; the line on the left is thin, darkcolored basal layer.
2. Specimen USNM Loc. $2387-11$ from the Cretaceous, Maastrichtian, near Maastricht (Limburg), Netherlands.
2a. Tangential thin-section $\times 30$. Shows section cutting two intrazoarial growth phases. Surface of juncture is about the middle of figure.
2b. Longitudinal thin-section $\times 30$. Shows parts of three overgrowth phases. Large cavity in proximal portion of middle unit is a broad chamber. In middle zoarial growth phase, intermediate diaphragms occur just aboral to terminal growth surface. Also, a few intermediate diaphragms are scattered throughout the growth unit.
Figure
Page
1, 2. Heteropora cryptopora (Goldfuss), 1826 ........................................ 115
Both specimens figured were collected from the Cretaceous, Maastrichtian, St. Petersberg near Maastricht (Limburg), Netherlands.
3. Paralectotype, UB 118b, figured by Goldfuss, $1826_{1}$ pl. 10, fig. 3b.
1a. Longitudinal section, acetate peel $\times 200$. Diaphragm in zooecial chamber on right shows slight aboral flexure and is interpreted as an intermediate diaphragm. Diaphragm in left-hand zooecium shows slight oral flexure at juncture with zooecial wall, but appearance may be due to recrystallization rather than primary structure. Zooecial chamber and zooecial wall to center right are continuous with suprajacent growth phase.
1 b. Longitudinal section, acetate peel $\times 200$. Shows distal portion of one growth unit and proximal portion of suprajacent intrazoarial overgrowth. Zooecial walls thin symmetrically near aperture. Note distinctly laminate basal layer of overgrowth, draping over apertural parts of zooecial wall. Poorly preserved intermediate diaphragm is shown in central zooecial chamber; diaphragm has short, aborally flexed abutments.
1c. Longitudinal section, acetate peel $\times 200$. Basal layer is undulatory, extends diagonally from upper right to lower left side of figure, and separates subjacent from suprajacent intrazoarial overgrowth (dark arrow). Intermediate diaphragms occur in zooecial chambers of subjacent growth unit (hollow arrows). Diaphragm in right center zooecium is dark in color and indistinctly laminate. Undulatory basal layer forms recumbent wall of most proximal overgrowth zooecium. More distal wall has optically structureless (originally granular?) calcareous tissue and thin zooecial lining.
4. Lectotype, UB 118a, was figured by Goldfuss, 1826, pl. 10, fig. 3a; by von Hagenow, 1851, pl. 5, fig. 6; and by Canu and Bassler, 1920, text fig. 222A, p. 681.
2a. Longitudinal section, acetate peel $\times 200$. Shows primary laminate structure of exozonal zooecial wall. Laminae are convex orally and continuous across zooecial boundary zone. Patches of homogeneous calcareous tissue occur in more aboral parts of outer cortex.
2 b. Longitudinal section, acetate peel $\times 500$. Zooecia grew orally to left. Shows apertural portion of zooecial wall, with zone of irregular structure at about center of picture making boundary between subjacent growth phase and suprajacent overgrowth phase. Zone of irregular structure is symmetrical with basal layer to either side. Zooecial chamber is continuous into overgrowth zooecium, but zooecial wall shows marked thinning. Also, note short stublike projection of zooecial wall in center part of figure (indicated by arrow), perhaps a remnant of a resorbed zooecial closure. The homogeneous appearance of cortex is probably secondary.





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## Explanation of Plate 36

Figure ..... Page

1. Leiosoecia parvicella (Gabb and Horn), 1860 ..... 123Lectotype, ANSP 31261, was probably figured by Gabb and Horn,1861, pl. 69, figs. 36-38, from the ?Paleocene, Vincentownformation, Timber Creek, New Jersey.
1a. Zoarial fragment $\times 2$ showing the intersection and anastomosis of distally growing branches.
1b. Surface of zoarium $\times 5$.
1c. Surface of zoarium $\times 30$.
1d. Transverse thin-section $\times 30$. Zooecia in endozone have polygonal outlines; in exozone, zoecial growth slightly undulatory. Zooecial walls have submoniliform to moniliform profiles.
1e. Longitudinal thin-section $\times 30$. Zooecial walls thin and parallel-sided to submoniliform in endozone, thickened and having moniliform profiles in exozone. Note slight asymmetry of thickness across zooecial boundary zone. Thin intermediate diaphragms seen in most zooecia.
1f. Tangential thin-section $\times 30$. Interzooidal pores uncommon. View shows no apparent arrangement of large and small polymorphs.
1 g. Longitudinal thin-section $\times 200$ showing microstructure, interzooidal pore, and thin intermediate diaphragms suboral to skeletal aperture.
1h. Tangential thin-section $\times 100$. Shows zooecial wall with amalgamate appearance and thin, dark zooecial linings.
Figure
2. Parleiosoecia jacksonica Canu and Bassler, 1920

Lectotype, USNM Loc. 2933B-1, previously figured by Canu and Bassler, 1920, pl. 148, fig. 2, from the Eocene, Jacksonian, Eutaw Springs, South Carolina.
1a. Zoarium $\times 5$. Zoarium has a large encrusting base and a single upright branch.
1b. Zoarial surface $\times 30$. Apertures of large polymorphs commonly have slightly raised rims. Each large polymorph is surrounded by numerous small polymorphs.
1c. Tangential thin-section $\times 30$. Note discontinuous variation in size between large and small polymorphs.
1d. Longitudinal thin-section at base of erect branch $\times 30$. Shows hollow axial chambers formed by distal extension of basal layer.
1e. Transverse thin-section $\times 30$. Shows hollow axial chamber, thin-walled polygonal zooecia in endozone, and thicker submoniliform walls in exozone.
1 f . Longitudinal thin-section $\times 100$. Shows cylindrical cortex structure composed of subgranular calcareous tissue.
1g. Tangential thin-section $\times 100$. Cortex of large polymorphs distinct in appearance from indistinctly laminated walls of small polymorphs.


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## Explanation of Plate 38

## Figure <br> Page

1-4. Parleiosoecia jacksonica Canu and Bassler, 1920
132
All specimens figured are topotypes, identified by R. S. Bassler, and collected from the Eocene, Jacksonian, Eutaw Springs, South Carolina.

1. USNM 2933A. Zoarial fragment $\times 5$ showing branching, and intersection and anastomosis of distally growing branches in upper right.
2. USNM 2933A-9. Longitudinal section, acetate peel $\times 5$ showing intersection and anastomosis of distally growing branches in upper right. The growing tip of the right-hand vertical branch abuts endozonal zooecia in the upper horizontal branch.
3. USNM Loc. 2933A-23.

3a. Longitudinal thin-section $\times 30$ showing intrazoarial overgrowth. Note that zooecial chamber in lower portion of figure is continuous with a zooecial chamber in the overgrowth; other encrusted zooecia are sealed orally by thin, intermediate (?) diaphragms.
3b. Longitudinal thin-section $\times 100$ from same branch as above showing profile and structure of zooecial walls. Note thin intermediate diaphragms in terminal position, and two intermediate diaphragms in zone of zooecial bend. Also note wall structure of large polymorph in upper part of figure showing distinct cortex structure, and the steeply dipping laminae in adjacent wall of small polymorph.
4. USNM 2933A-26.

4a. Tangential section, acetate peel $\times 300$ showing wall structure of large polymorphs and small polymorphs. Small polymorphs distinctly laminate; cortex of large polymorphs subgranular.
4b. Longitudinal thin-section $\times 30$ showing axial chambers formed by basal layer and the intersection and anastomosis of two separate branches. A dark line marks surface of intersection in central and inner peripheral area but is lost in outer peripheral area. Note nearly $180^{\circ}$ reflection of zooecial growth axes in lower part of figure so that zooecia grow proximally in relation to growth of branches.
$4 c$. Detail of $1 \mathrm{~b} \times 100$ showing wedging out of zooecial cavities at surface of intersection. Zooecial walls merge continuously at surface of intersection to form a thin common wall.

## Explanation of Plate 39

Figure
1-5. Parleiosoecia jacksonica Canu and Bassler, 1920 ..... 132

Figs. 1, 3-5 are specimens from the Eocene, Jacksonian, Eutaw Springs, South Carolina; fig. 2, specimens from Eocene, Jacksonian, Santee River, three miles above Lenuds Ferry, South Carolina.

1. USNM Loc. 2933A, branch $\times 5$.
2. USNM Cat. 65447-1, figured by Canu and Bassler, 1922, pl. 148 , fig. 1.
2a. Surface of branch $\times 30$. Shows apertures of large polymorphs with slightly raised rims surrounded by apertures of small polymorphs. Note regular, nearly rhombohedral, arrangements of large polymorphs.
2b. Longitudinal thin-section $\times 100$ showing axial chambers formed by basal layer. Note dark line separating basal layer from recumbent zooecial wall, and intermediate diaphragms in zone of zooecial bend.
2c. Detail of $2 \mathrm{~b} \times 300$ showing laminated structure of basal layer.
3. USNM Loc. 2933A-14. Longitudinal section, acetate peel $\times 5$ showing characteristic appearance of branch axis and coaxial exozone-endozone.
4. USNM Loc. 2933A-26. Longitudinal thin-section $\times 100$ showing axial chambers formed by basal layer. Note distal flexure of basal layer in center.
5. USNM Loc. 2933A-23. Transverse thin-section $\times 100$ showing basal layer and shape and arrangement of zooecial openings.



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## Explanation of Plate 40

Figure

1. Parleiosoecia iacksonica Canu and Bassler, 1920 ...................... 132
USNM Cat. 65449-1, Eocene, middle Jacksonian, 18 miles west
of Wrightsville, Georgia. Specimen figured by Canu and
Bassler, 1922, pl. 148, fig. 6 .

## Explanation of Plate 41

Figure ..... Page

1. Reptonodicava globosa (Michelin), 1846 ..... 140MNHN d’Orb. Coll. 2988-1, Jurassic, Bathonien, Luc (Calvados),France.
1a. Clavate zoarium $\times 2.5$. Surface is smooth.
1b. Surface of zoarial fragment $\times 30$.
1c. Longitudinal thin-section $\times 5$. Faint, dark-colored bands symmetrical with zoarial surface are probably abandoned growth surfaces. Outermost living chambers are closed aborally by basal diaphragms, and chambers are filled with dark, fine-grained sediments. Interior zooecial cavities are filled by clear calcite.
1d. Tangential thin-section $\times 30$.
1e. Longitudinal thin-section $\times 100$ showing thin, dark, basal diaphragms; diaphragms flex orally and form abutments.
1f. Longitudinal thin-section, crossed nicols $\times 400$. Laminae appear to be oriented orally acute from boundary zone, then arch broadly across the cortex and are directed aborally in the inner cortex.
1 g . Longitudinal thin-section $\times 30$. Shows submoniliform to moniliform zooecial walls; repetition of dark bands formed by a thin, dark zone with zooecial wall (detail of 1c) suggests cyclic growth intervals.
1h. Transverse thin-section $\times 30$ showing moniliform profile of zooecial walls and basa! diaphragms.

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## Explanation of Plate 42

Figure ..... Page

1. Reptonodicava globosa (Michelin), 1846 ..... 140MNHN d'Orb. Coll. 2988-2, Jurassic, Bathonien, Luc (Calvados), France.
1a. Zoarium $\times 2.5$. Zoarium is a hemispherical mass raised slightly above the substrate on narrower peduncle-like base.
1b. Surface of zoarium $\times 30$.
1c. Longitudinal thin-section $\times 5$. Intrazoarial overgrowth is seen in the proximal portion of colony.
1d. Longitudinal thin-section $\times 20$. Zooecial walls have submoniliform to moniliform profiles, walls typically equal and symmetrical in thickness across the boundary zone.
1e. Tangential thin-section $\times 30$.
1f. Longitudinal thin-section $\times 100$. Shows essentially continuously growing zooecial wall on right, and zooecial walls budding from basal layer on left. Zooecial wall, growing continuously, shows little variation in thickness. Zooecial walls newly budded from basal layer are very thin initially, but attain normal exozonal thickness and appearance within a very short distance.
1g. Longitudinal thin-section $\times 100$. Zooecia grew towards upper right. Walls of encrusting zooecia are continuous with thin, basal layer. Terminal diaphragms of subjacent zooecia are separated from encrusting basal layer by a dark line.

## Explanation of Plate 43

Figure ..... Page
1-3. Reptonodicava globosa (Michelin), 1846 ..... 140All specimens from Jurassic, Bathonien, Luc (Calvados), France.1. MNHN d'Orb. Coll. 2988-1. Longitudinal thin-section $\times 400$showing basal diaphragm. Diaphragm flexes and extendsorally as a thin abutment. Zooecial walls nearly same coloras clear calcite filling zooecial chamber, but show greaterdensity of small dark granules.
2. MNHN d'Orb. Coll. 2988-2, tangential thin-section $\times 100$. Numerous, short, blunt, mural spines give slightly crenulate appearance to outline of zooecial cavity. Note fine laminations in outer cortex and zooecial lining.
3. MNHN d'Orb. Coll. 2988-3.
3a. Longitudinal thin-section $\times 5$. Shows typical growth habit and local intrazoarial overgrowth in upper left portion of zoarium. Fine-grained, dark-colored sediment fills lastformed living chambers and portions of zooecial cavities just subjacent to local intrazoarial overgrowth.
3b. Longitudinal thin-section $\times 30$, detail of local intrazoarial encrustation shown in 3a. Outer zooecial chamber subjacent to overgrowth is closed aborally by intermediate diaphragms and filled with dark, fine-grained sediment.
3c. Longitudinal thin-section $\times 100$. Shows zooecial walls with moniliform profile and interzooidal pores. Thin, basal diaphragms are poorly preserved. Zooecial walls show obscure lamination.
3d. Longitudinal thin-section $\times 100$. Zooecia grew towards upper right. Note diaphragm subjacent to basal layer of intrazoarial encrustation. Diaphragm shows slight aboral flexure at juncture with zooecial wall and is apparently continuous with it.



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## Explanation of Plate 44

Figure Page
1,2. Repłonodicava globosa (Michelin), 1846 ..... 140
Both specimens from Jurassic, Bathonien, Ranville (Calvados), France.

1. USNM 32171-4, longitudinal section $\times 5$. Section includes encrusting basal portion.
2. USNM 32171-1.
2a. Longitudinal thin-section $\times 5$, branch from a more massiveappearing zoarium. Dark bands suggest periodic addition of growth increments. Note numerous, evenly spaced basal diaphragms and very long, relatively straight zooccial located near major vector of branch growth.
2b. Longitudinal thin-section $\times 5$; another branchlike extension from the same zoarium. Various stages in zoarial growth are shown by dark bands, indicating that once budded zooecia continued to grow for most of the life of the colony, but grew at different rates. This part of the zoarium is overgrown by an encrusting cyclostome, c.f. Berenicea.
2c. Longitudinal thin-section $\times 20$, detail of 2 a . Appearance of banding separating growth increments is partly due to inclusion of dark material in walls, gradual increase in wall thickness in a given increment continued orally by reduced thickness, and regular spacing of diaphragms. Major increments apparently include many minor incremental episodes.
Figure Page
3. Tełrocycloecia dichołoma Canu, 1917 ..... 148NMW 1859 L686-1, misidentified by Reuss as Heteropora dicho-toma (Goldfuss), from Miocene, Tortonian, Leithakalke, Eisen-stadt, Hungary.

1a. Branch $\times 5$ showing regular distribution of apertures of large polymorphs.
1b. Tangential thin-section $\times 30$ showing discontinuous variation in size between openings of large and small polymorphs.
1c. Surface of zoarium $\times 30$. Surface is slightly worn. Note light-colored rings of calcareous tissue around apertures, apparently reflecting microstructure of zooecial wall; e.g. compare to 1 e .
1d. Transverse thin-section $\times 30$. Exozone and endozone broadly gradational; zooecial walls in exozone submoniliform and slightly undulatory.
1e. Tangential thin-section $\times 100$. Zooecial walls in this view have integrate appearance because of concentration of dark granules along boundary zone; walls also have very thin, dark, laminate zooecial linings.
1f. Longitudinal thin-section $\times 30$. Note relatively simple appearance. Exozone and endozone are gradational with broad zooecial curve from endozone; zooecia typically intersect zoarial surface at less than $90^{\circ}$. Zooecial walls display submoniliform profiles in both endozone and exozone.
1 g . Longitudinal thin-section $\times 100$, detail of 1 f . Shows zooecial wall with indistinct, orally convex laminae in cortex, and thin, dark, zooecial lining.


1 a


1 b


1 d


1 e



Cerioporid Cyclostomes (Bryozoa): Nye 215

## Explanation of Plate 46

Figure Page
1-3. Tetrocycloecia dichotoma Canu, 1918 ..... 148
All specimens figured were misidentified by Reuss as Heteroporadichotoma (Goldfuss), and were from the Miocene, Tortonian,Leithekalke, Eisenstadt, Hungary.

1. NMW 1859 L686-2.
1a. Branch $\times 5$.
1b. Longitudinal thin-section $\times 30$. Zooecial walls submoniliformin both endozone and exozone.
1c. Tangential thin-section $\times 30$.
1d. Transverse thin-section $\times 30$.
2. NMW 1867-1, previously figured by Manzoni, 1877, pl. 12,fig. 46 as Heteropora dichotoma (Goldfuss).
2a. Branch $\times 5$.
2b. Longitudinal thin-section $\times 30$. Zooecia grow oblique frommajor vector of branch growth.
3. NMW 1859 L686-1 $\times 30$. Shows unworn portion of zoarial surface in lower right, and apertures of large polymorphs with slightly raised prominent rims.

Explanation of Plate 47
Figure Page

1. Zonopora spiralis (Goldfuss), 1826 ..... 156Lectotype UB 133. Specimen is probably the specimen figured byGoldfuss, 1826, pl. 11, figs. 2a, b, from Cretaceous, Maastrich-tian, St. Petersberg near Maastricht (Limburg), Netherlands.
1a. Branch $\times 5$ showing typical helical appearance of zoaria. The convexly curved zoarial surface is referred to as the zoarial salient; the concave surface as the zoarial embayment. Terminal diaphragms appear as a nearly continuous sheet over the upper portion of the embayment.
1b. Surface of branch $\times 20$, detail of 1 a .
1c. Deep tangential thin-section $\times 30$.
1d. Tangential thin-section $\times 100$. Deep tangential section of large zooecia showing amalgamate appearance and thin, laminate, zooecial linings.
1e. Transverse thin-section $\times 30$. Large zooecia on lefthand side opening at zoarial salient, small zooecia on right opening at zoarial embayment. Branch axis is offset to right center.
1f. Longitudinal thin-section $\times 30$. Large zooecia directed towards zoarial salient on right, small zooecia towards zoarial embayment on left. Note terminal diaphragm in small zooecia at left.
1 g . Tangential thin-section $\times 30$; relatively shallow section of large zooecia showing thick zooecial linings. Note spinelike projections of homogeneous-appearing cortex tissue towards zooecial chamber. Mural spines are commonly submerged by thick zooecial lining in outer exozone.


1 a


1 b

l c


1 d





1 d


## Explanation of Plate 48

Figure Page

1. Zonopora spiralis (Goldfuss), 1826 ..... 156USNM Loc. 2965A-25, Cretaceous, Mastrichtian, Mastricht(Limburg), Netherlands.
1a. Longitudinal section, acetate peel $\times 5$. Shows profile with zoarial salients and embayments resulting from helical growth about a major vector of distal growth. Large zooecia intersect surface of salient at nearly right angles; small zooecia intersect surface of embayment obliquely. Branch to lower left appears to be an intrazoarial overgrowth.
1 b . Tangential section, acetate peel $\times 30$. Shallow tangential section of small zooecia, deep tangential section of large zooecia.
1c. Longitudinal section, acetate peel $\times 30$, detail from 1a. Zooecial walls in endozone thin and parallel-sided. Exozonal zooecial walls thicker, sometimes submoniliform, and have characteristic lanceolate profile.
1d. Longitudinal section, acetate peel $\times 100$, detail of 1 c showing typical wall structure with light-colored cortex tissue bounded by thick, dark-colored deposits of zooecial lining. Spinelike lateral projections of cortex tissue submerged by zooecial lining, but sometimes project into zooecial cavity as mural spines.
1e. Longitudinal section, acetate peel $\times 100$ showing wall structure of small zooecia and interzooidal pores. Note thick deposits of dark-colored, longitudinally laminated tissue.
1f. Tangential section, acetate peel $\times 100$. Section is just distal to 1 e , showing shallow tangential view of small zooecia. Note amalgamate, granular appearance of zooecial walls and thick zooecial linings.
1 g . Tangential section, acetate peel $\times 100$. Section is moderately deep, showing thin zooecial lining. Note mural spines and narrow, straight, interzooidal pore in lower left quadrant of figure.

## Explanation of Plate 49

Figure Page

1. Zonopora spiralis (Goldfuss), 1826 ..... 156USNM Loc. 2965A-7, Cretaceous, Maastrichtian, Maastricht(Limburg), Netherlands.
1a. Longitudinal section, acetate peel $\times 30$.1b. Transverse section, acetate peel $\times 30$. Large zooecia onleft, small zooecia on right. Axis of branch is shifted toright of center; note thick zooecial linings in exozone.
1c. Tangential section, acetate peel $\times 100$, detail of large zooecia in 1 b . Note increased thickness of zooecial lining in exozone.
1d. Tangential section, acetate peel $\times 30$. Upper part of figure shows shallow tangential section of small zooecia of zoarial embayment and deeper tangential view of large zooecia in zoarial salient.
1e. Longitudinal section, acetate peel $\times 100$. The zooecial boundary zone is marked by a narrow zone of light-colored, homogeneous tissue bounded by the indistinctly laminated cortex tissue. Lamination in the cortex, as seen in second zooecial wall from right, is only slightly curved convex orally, and abuts the zooecial lining at $60^{\circ}-70^{\circ}$. Zooecial linings are characteristically thick and dark in color. A narrow, straight, interzooidal pore is seen in aboral part of second zooecial wall from right. Mural spines extend into zooecial cavity in more aboral portion of zooecial wall in left part of figure. Spinelike extensions from cortex are submerged by thick zooecial linings in zooecial walls in righthand portion of figure.


1 a




1 b


## Explanation of Plate 50

Figure ..... Page
1-3. Zonopora spiralis (Goldfuss), 1826 ..... 156All specimens from Cretaceous, Maastrichtian, Maastricht (Lim-burg), Netherlands.

1. USNM Loc. 2965A-21.
1a. Longitudinal section, acetate peel $\times 30$ showing small zooecia at zoarial embayment with porous terminal diaphragms. Primary growth is encrusted by intrazoarial overgrowth.
1b. Longitudinal section, acetate peel $\times 100$, detail of 1 a . Terminal diaphragm is porous and similar in appearance to zooecial lining of subjacent zooecial walls. Adjacent diaphragms are not continuous across terminus of zooecial wall. Light-colored, homogeneous tissue of basal layer is continuous with cortex tissue of encrusting zooecia.
2. USNM Loc. $2965 \mathrm{~A}-23$.
2a. Longitudinal section, acetate peel $\times 30$. Shows large zooecia budding from axial region offset to right side of branch. Shows thin, parallel-sided walls in endozone and thickened exozonal walls with characteristic club or lanceolate profiles. Note that the apertural, rims are constricted, forming a cusp-shaped profile from which terminal diaphragms extend laterally.
2b. Longitudinal section, acetate peel $\times 100$. Tissue of terminal diaphragm is continuous with zooecial linings, but does not extend across light-colored cortex tissue at terminus of zooecial wall.
2c. Longitudinal section, acetate peel $\times 100$. Detail of 1 a showing cuspate extension of apertural rim and lateral flexure of zooecial lining to form terminal diaphragm.
3. USNM Loc. 2965A-26, longitudinal section, acetate peel $\times 100$. Zooecial walls of large zooecia in outer exozone. The thin zooecial lining and rodlike extension of cortex into the zooecial cavity as mural spines, suggest a younger ontogenetic stage than shown in Pl. 36, figs. 1d, e, and pl. 37, fig. 1 e.

Explanation of Plate 51

Figure
Page

1. Recent cerioporid, USNM $6086-1, \times 30$. Brood chamber with roof partially broken away to show interior, revealing intrachamber zooecia connected by thin septate walls.
2. Recent cerioporid, BM O'Donoghue Coll. 1963-2-6-1 pt. all $\times 100$. Soft tissues are stained.
2a. Tangential thin-section. Basal portion of tentacular crowns in three zooecia; brown bodies in smaller zooecia to left. Dark lines in skeletal wal! probably are algal or fungal borings. Note dark staining nuclei of cellular tissue in interzooidal part indicated by arrow.
2b. Longitudinal thin-section. Shows tentacular crowns and visceral sacs of two zooids. Insertion of lophophore retractor muscles directly on body wall shown in zooid on right; note lack of funiculus at base of visceral sac.
2c. Shows tentacular crown, visceral sac and brown bodies in center zooecium. Nucleated strands of tissue appear to pass continuously through interzooidal pore marked by arrow.


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    1. Lectotype UB 120 ; and one identified specimen from near the type locality, USNM 2405-2.
[^4]:    I think, however, that von Hagenow's ideas as to the suitable limits of a species were a little exaggerated, and I am not convinced that the specimens figured by Goldfuss (op. cit.) in his Pl. X, figs. 3a, 3b, and 3c, did not all belong to one and the same species. The differences existing between them seem to me to be easily explainable by the fact that they obviously represent different portions of three zoaria.

[^5]:    1920. Pars Leiosoecia Canu, and Bassler, U.S. Nat. Mus., Bull. 106, p. 823.
    1921. Pars Leiosoecia Canu and Bassler, Canu, and Bassler, U.S. Nat. Mus., Proc., vol. 61, p. 99.
[^6]:    *In millimeters

[^7]:    *In millimeters
    Exozone 1: Outer exozone; measurements made from tangential section.

[^8]:    KEY TO SPECIMEN CODE

    1. Topotype specimens identified by Reuss (Syntype?) : NMW 1859 L686-1, 2; NMW 1867xL1-1.
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    3. NMW 1859 L686-1
    4. NMW 1859 L686-1 ( 8$)$; NMW 1867xL1-1 (7).
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